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(54) Title: NUCLEIC ACIDS AND PROTEINS FROM STREPTOCOCCUS GROUPS A & B

(57) Abstract: The invention provides proteins from group B streptococcus (*Streptococcus agalactiae*) and group A streptococcus (*Streptococcus pyogenes*), including amino acid sequences and the corresponding nucleotide sequences. Data are given to show that the proteins are useful antigens for vaccines, immunogenic compositions, and/or diagnostics. The proteins are also targets for antibiotics.

NUCLEIC ACIDS AND PROTEINS FROM STREPTOCOCCUS GROUPS A & B

All documents cited herein are incorporated by reference in their entirety.

TECHNICAL FIELD

This invention relates to nucleic acid and proteins from the bacteria *Streptococcus agalactiae* (GBS) and
5 *Streptococcus pyogenes* (GAS).

BACKGROUND ART

Once thought to infect only cows, the Gram-positive bacterium *Streptococcus agalactiae* (or "group B streptococcus", abbreviated to "GBS") is now known to cause serious disease, bacteremia and meningitis, in immunocompromised individuals and in neonates. There are two types of neonatal
10 infection. The first (early onset, usually within 5 days of birth) is manifested by bacteremia and pneumonia. It is contracted vertically as a baby passes through the birth canal. GBS colonises the vagina of about 25% of young women, and approximately 1% of infants born via a vaginal birth to colonised mothers will become infected. Mortality is between 50-70%. The second is a meningitis that occurs 10 to 60 days after birth. If pregnant women are vaccinated with type III capsule so that the infants are
15 passively immunised, the incidence of the late onset meningitis is reduced but is not entirely eliminated.

The "B" in "GBS" refers to the Lancefield classification, which is based on the antigenicity of a carbohydrate which is soluble in dilute acid and called the C carbohydrate. Lancefield identified 13 types of C carbohydrate, designated A to O, that could be serologically differentiated. The organisms that most commonly infect humans are found in groups A, B, D, and G. Within group B, strains can be
20 divided into 8 serotypes (Ia, Ib, Ia/c, II, III, IV, V, and VI) based on the structure of their polysaccharide capsule.

Group A streptococcus ("GAS", *S.pyogenes*) is a frequent human pathogen, estimated to be present in between 5-15% of normal individuals without signs of disease. When host defences are compromised, or when the organism is able to exert its virulence, or when it is introduced to vulnerable tissues or hosts,
25 however, an acute infection occurs. Diseases include puerperal fever, scarlet fever, erysipelas, pharyngitis, impetigo, necrotising fasciitis, myositis and streptococcal toxic shock syndrome.

S.pyogenes is typically treated using antibiotics. Although *S.agalactiae* is inhibited by antibiotics, however, it is not killed by penicillin as easily as GAS. Prophylactic vaccination is thus preferable.

Current GBS vaccines are based on polysaccharide antigens, although these suffer from poor
30 immunogenicity. Anti-idiotypic approaches have also been used (e.g. WO99/54457). There remains a need, however, for effective adult vaccines against *S.agalactiae* infection. There also remains a need for vaccines against *S.pyogenes* infection.

It is an object of the invention to provide proteins which can be used in the development of such vaccines. The proteins may also be useful for diagnostic purposes, and as targets for antibiotics.

DISCLOSURE OF THE INVENTION

The invention provides proteins comprising the *S.agalactiae* amino acid sequences disclosed in the examples, and proteins comprising the *S.pyogenes* amino acid sequences disclosed in the examples. These amino acid sequences are the even SEQ IDs between 1 and 10960.

- 5 It also provides proteins comprising amino acid sequences having sequence identity to the *S.agalactiae* amino acid sequences disclosed in the examples, and proteins comprising amino acid sequences having sequence identity to the *S.pyogenes* amino acid sequences disclosed in the examples. Depending on the particular sequence, the degree of sequence identity is preferably greater than 50% (e.g. 60%, 70%, 80%, 90%, 95%, 99% or more). These proteins include homologs, orthologs, allelic variants and
- 10 functional mutants. Typically, 50% identity or more between two proteins is considered to be an indication of functional equivalence. Identity between proteins is preferably determined by the Smith-Waterman homology search algorithm as implemented in the MPSRCH program (Oxford Molecular), using an affine gap search with parameters *gap open penalty*=12 and *gap extension penalty*=1.

- 15 Preferred proteins of the invention are GBS1 to GBS689 (see Table IV).

The invention further provides proteins comprising fragments of the *S.agalactiae* amino acid sequences disclosed in the examples, and proteins comprising fragments of the *S.pyogenes* amino acid sequences disclosed in the examples. The fragments should comprise at least *n* consecutive amino acids from the sequences and, depending on the particular sequence, *n* is 7 or more (e.g. 8, 10, 12, 14, 16, 18, 20, 30,

20 40, 50, 60, 70, 80, 90, 100 or more). Preferably the fragments comprise one or more epitopes from the sequence. Other preferred fragments are (a) the N-terminal signal peptides of the proteins disclosed in the examples, (b) the proteins disclosed in the examples, but without their N-terminal signal peptides, (c) fragments common to the related GAS and GBS proteins disclosed in the examples, and (d) the proteins disclosed in the examples, but without their N-terminal amino acid residue.

- 25 The proteins of the invention can, of course, be prepared by various means (e.g. recombinant expression, purification from GAS or GBS, chemical synthesis *etc.*) and in various forms (e.g. native, fusions, glycosylated, non-glycosylated *etc.*). They are preferably prepared in substantially pure form (i.e. substantially free from other streptococcal or host cell proteins) or substantially isolated form. Proteins of the invention are preferably streptococcal proteins.

- 30 According to a further aspect, the invention provides antibodies which bind to these proteins. These may be polyclonal or monoclonal and may be produced by any suitable means (e.g. by recombinant expression). To increase compatibility with the human immune system, the antibodies may be chimeric or humanised (e.g. Breedveld (2000) *Lancet* 355(9205):735-740; Gorman & Clark (1990) *Semin. Immunol.* 2:457-466), or fully human antibodies may be used. The antibodies may include a detectable label (e.g. for diagnostic assays).
- 35

According to a further aspect, the invention provides nucleic acid comprising the *S.agalactiae* nucleotide sequences disclosed in the examples, and nucleic acid comprising the *S.pyogenes* nucleotide sequences disclosed in the examples. These nucleic acid sequences are the odd SEQ IDs between 1 and 10966.

5 In addition, the invention provides nucleic acid comprising nucleotide sequences having sequence identity to the *S.agalactiae* nucleotide sequences disclosed in the examples, and nucleic acid comprising nucleotide sequences having sequence identity to the *S.pyogenes* nucleotide sequences disclosed in the examples. Identity between sequences is preferably determined by the Smith-Waterman homology search algorithm as described above.

10 Furthermore, the invention provides nucleic acid which can hybridise to the *S.agalactiae* nucleic acid disclosed in the examples, and nucleic acid which can hybridise to the *S.pyogenes* nucleic acid disclosed in the examples preferably under 'high stringency' conditions (e.g. 65°C in 0.1xSSC, 0.5% SDS solution).

Nucleic acid comprising fragments of these sequences are also provided. These should comprise at least
15 *n* consecutive nucleotides from the *S.agalactiae* or *S.pyogenes* sequences and, depending on the particular sequence, *n* is 10 or more (e.g. 12, 14, 15, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200 or more). The fragments may comprise sequences which are common to the related GAS and GBS sequences disclosed in the examples.

According to a further aspect, the invention provides nucleic acid encoding the proteins and protein
20 fragments of the invention.

The invention also provides: nucleic acid comprising nucleotide sequence SEQ ID 10967; nucleic acid comprising nucleotide sequences having sequence identity to SEQ ID 10967; nucleic acid which can hybridise to SEQ ID 10967 (preferably under 'high stringency' conditions); nucleic acid comprising a fragment of at least *n* consecutive nucleotides from SEQ ID 10967, wherein *n* is 10 or more e.g. 12, 14,
25 15, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250, 300, 350, 400, 450, 500, 600, 700, 800, 900, 1000, 1500, 2000, 3000, 4000, 5000, 10000, 100000, 1000000 or more

Nucleic acids of the invention can be used in hybridisation reactions (e.g. Northern or Southern blots, or in nucleic acid microarrays or 'gene chips') and amplification reactions (e.g. PCR, SDA, SSSR, LCR, TMA, NASBA *etc.*) and other nucleic acid techniques.

30 It should also be appreciated that the invention provides nucleic acid comprising sequences complementary to those described above (e.g. for antisense or probing, or for use as primers).

Nucleic acid according to the invention can, of course, be prepared in many ways (e.g. by chemical synthesis, from genomic or cDNA libraries, from the organism itself *etc.*) and can take various forms (e.g. single stranded, double stranded, vectors, primers, probes, labelled *etc.*). The nucleic acid is
35 preferably in substantially isolated form.

Nucleic acid according to the invention may be labelled *e.g.* with a radioactive or fluorescent label. This is particularly useful where the nucleic acid is to be used in nucleic acid detection techniques *e.g.* where the nucleic acid is a primer or as a probe for use in techniques such as PCR, LCR, TMA, NASBA *etc.*

5 In addition, the term "nucleic acid" includes DNA and RNA, and also their analogues, such as those containing modified backbones, and also peptide nucleic acids (PNA) *etc.*

According to a further aspect, the invention provides vectors comprising nucleotide sequences of the invention (*e.g.* cloning or expression vectors) and host cells transformed with such vectors.

10 According to a further aspect, the invention provides compositions comprising protein, antibody, and/or nucleic acid according to the invention. These compositions may be suitable as immunogenic compositions, for instance, or as diagnostic reagents, or as vaccines.

The invention also provides nucleic acid, protein, or antibody according to the invention for use as medicaments (*e.g.* as immunogenic compositions or as vaccines) or as diagnostic reagents. It also provides the use of nucleic acid, protein, or antibody according to the invention in the manufacture of: (i) a medicament for treating or preventing disease and/or infection caused by streptococcus; (ii) a
15 diagnostic reagent for detecting the presence of streptococcus or of antibodies raised against streptococcus; and/or (iii) a reagent which can raise antibodies against streptococcus. Said streptococcus may be any species, group or strain, but is preferably *S.agalactiae*, especially serotype III or V, or *S.pyogenes*. Said disease may be bacteremia, meningitis, puerperal fever, scarlet fever, erysipelas, pharyngitis, impetigo, necrotising fasciitis, myositis or toxic shock syndrome.

20 The invention also provides a method of treating a patient, comprising administering to the patient a therapeutically effective amount of nucleic acid, protein, and/or antibody of the invention. The patient may either be at risk from the disease themselves or may be a pregnant woman ('maternal immunisation' *e.g.* Glezen & Alpers (1999) *Clin. Infect. Dis.* 28:219-224).

Administration of protein antigens is a preferred method of treatment for inducing immunity.

25 Administration of antibodies of the invention is another preferred method of treatment. This method of passive immunisation is particularly useful for newborn children or for pregnant women. This method will typically use monoclonal antibodies, which will be humanised or fully human.

The invention also provides a kit comprising primers (*e.g.* PCR primers) for amplifying a template sequence contained within a *Streptococcus* (*e.g.* *S.pyogenes* or *S.agalactiae*) nucleic acid sequence, the
30 kit comprising a first primer and a second primer, wherein the first primer is substantially complementary to said template sequence and the second primer is substantially complementary to a complement of said template sequence, wherein the parts of said primers which have substantial complementarity define the termini of the template sequence to be amplified. The first primer and/or the second primer may include a detectable label (*e.g.* a fluorescent label).

The invention also provides a kit comprising first and second single-stranded oligonucleotides which allow amplification of a *Streptococcus* template nucleic acid sequence contained in a single- or double-stranded nucleic acid (or mixture thereof), wherein: (a) the first oligonucleotide comprises a primer sequence which is substantially complementary to said template nucleic acid sequence; (b) the second oligonucleotide comprises a primer sequence which is substantially complementary to the complement of said template nucleic acid sequence; (c) the first oligonucleotide and/or the second oligonucleotide comprise(s) sequence which is not complementary to said template nucleic acid; and (d) said primer sequences define the termini of the template sequence to be amplified. The non-complementary sequence(s) of feature (c) are preferably upstream of (i.e. 5' to) the primer sequences. One or both of these (c) sequences may comprise a restriction site (e.g. EP-B-0509612) or a promoter sequence (e.g. EP-B-0505012). The first oligonucleotide and/or the second oligonucleotide may include a detectable label (e.g. a fluorescent label).

The template sequence may be any part of a genome sequence (e.g. SEQ ID 10967). For example, it could be a rRNA gene (e.g. Turenne *et al.* (2000) *J. Clin. Microbiol.* 38:513-520; SEQ IDs 12018-12024 herein) or a protein-coding gene. The template sequence is preferably specific to GBS.

The invention also provides a computer-readable medium (e.g. a floppy disk, a hard disk, a CD-ROM, a DVD *etc.*) and/or a computer database containing one or more of the sequences in the sequence listing. The medium preferably contains SEQ ID 10967.

The invention also provides a hybrid protein represented by the formula $\text{NH}_2\text{-A-}[\text{X-L}]_n\text{-B-COOH}$, wherein X is a protein of the invention, L is an optional linker amino acid sequence, A is an optional N-terminal amino acid sequence, B is an optional C-terminal amino acid sequence, and n is an integer greater than 1. The value of n is between 2 and x , and the value of x is typically 3, 4, 5, 6, 7, 8, 9 or 10. Preferably n is 2, 3 or 4; it is more preferably 2 or 3; most preferably, $n = 2$. For each n instances, -X- may be the same or different. For each n instances of $[\text{X-L}]$, linker amino acid sequence -L- may be present or absent. For instance, when $n=2$ the hybrid may be $\text{NH}_2\text{-X}_1\text{-L}_1\text{-X}_2\text{-L}_2\text{-COOH}$, $\text{NH}_2\text{-X}_1\text{-X}_2\text{-COOH}$, $\text{NH}_2\text{-X}_1\text{-L}_1\text{-X}_2\text{-COOH}$, $\text{NH}_2\text{-X}_1\text{-X}_2\text{-L}_2\text{-COOH}$, *etc.* Linker amino acid sequence(s) -L- will typically be short (e.g. 20 or fewer amino acids i.e. 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples include short peptide sequences which facilitate cloning, poly-glycine linkers (i.e. Gly $_n$ where $n = 2, 3, 4, 5, 6, 7, 8, 9, 10$ or more), and histidine tags (i.e. His $_n$ where $n = 3, 4, 5, 6, 7, 8, 9, 10$ or more). Other suitable linker amino acid sequences will be apparent to those skilled in the art. -A- and -B- are optional sequences which will typically be short (e.g. 40 or fewer amino acids i.e. 39, 38, 37, 36, 35, 34, 33, 32, 31, 30, 29, 28, 27, 26, 25, 24, 23, 22, 21, 20, 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples include leader sequences to direct protein trafficking, or short peptide sequences which facilitate cloning or purification (e.g. histidine tags i.e. His $_n$ where $n = 3, 4, 5, 6, 7, 8, 9, 10$ or more). Other suitable N-terminal and C-terminal amino acid sequences will be apparent to those

skilled in the art. In some embodiments, each X will be a GBS sequence; in others, mixtures of GAS and GBS will be used.

According to further aspects, the invention provides various processes.

5 A process for producing proteins of the invention is provided, comprising the step of culturing a host cell of to the invention under conditions which induce protein expression.

A process for producing protein or nucleic acid of the invention is provided, wherein the protein or nucleic acid is synthesised in part or in whole using chemical means.

10 A process for detecting polynucleotides of the invention is provided, comprising the steps of: (a) contacting a nucleic probe according to the invention with a biological sample under hybridising conditions to form duplexes; and (b) detecting said duplexes.

A process for detecting *Streptococcus* in a biological sample (e.g. blood) is also provided, comprising the step of contacting nucleic acid according to the invention with the biological sample under hybridising conditions. The process may involve nucleic acid amplification (e.g. PCR, SDA, SSSR, LCR, TMA, NASBA etc.) or hybridisation (e.g. microarrays, blots, hybridisation with a probe in solution etc.). PCR detection of *Streptococcus* in clinical samples, in particular *S.pyogenes*, has been reported [see e.g. Louie et al. (2000) *CMAJ* 163:301-309; Louie et al. (1998) *J. Clin. Microbiol.* 36:1769-1771]. Clinical assays based on nucleic acid are described in general in Tang et al. (1997) *Clin. Chem.* 43:2021-2038.

20 A process for detecting proteins of the invention is provided, comprising the steps of: (a) contacting an antibody of the invention with a biological sample under conditions suitable for the formation of an antibody-antigen complexes; and (b) detecting said complexes.

A process for identifying an amino acid sequence is provided, comprising the step of searching for putative open reading frames or protein-coding regions within a genome sequence of *S.agalactiae*. This will typically involve *in silico* searching the sequence for an initiation codon and for an in-frame termination codon in the downstream sequence. The region between these initiation and termination codons is a putative protein-coding sequence. Typically, all six possible reading frames will be searched. Suitable software for such analysis includes ORFFINDER (NCBI), GENEMARK [Borodovsky & McIninch (1993) *Computers Chem.* 17:122-133], GLIMMER [Salzberg et al. (1998) *Nucleic Acids Res.* 26:544-548; Salzberg et al. (1999) *Genomics* 59:24-31; Delcher et al. (1999) *Nucleic Acids Res.* 27:4636-4641], or other software which uses Markov models [e.g. Shmatkov et al. (1999) *Bioinformatics* 15:874-876]. The invention also provides a protein comprising the identified amino acid sequence. These proteins can then expressed using conventional techniques.

35 The invention also provides a process for determining whether a test compound binds to a protein of the invention. If a test compound binds to a protein of the invention and this binding inhibits the life cycle of the GBS bacterium, then the test compound can be used as an antibiotic or as a lead compound for the

design of antibiotics. The process will typically comprise the steps of contacting a test compound with a protein of the invention, and determining whether the test compound binds to said protein. Preferred proteins of the invention for use in these processes are enzymes (e.g. tRNA synthetases), membrane transporters and ribosomal proteins. Suitable test compounds include proteins, polypeptides, carbohydrates, lipids, nucleic acids (e.g. DNA, RNA, and modified forms thereof), as well as small organic compounds (e.g. MW between 200 and 2000 Da). The test compounds may be provided individually, but will typically be part of a library (e.g. a combinatorial library). Methods for detecting a binding interaction include NMR, filter-binding assays, gel-retardation assays, displacement assays, surface plasmon resonance, reverse two-hybrid *etc.* A compound which binds to a protein of the invention can be tested for antibiotic activity by contacting the compound with GBS bacteria and then monitoring for inhibition of growth. The invention also provides a compound identified using these methods.

The invention also provides a composition comprising a protein of the invention and one or more of the following antigens:

- 15 – a protein antigen from *Helicobacter pylori* such as VacA, CagA, NAP, HopX, HopY [e.g. WO98/04702] and/or urease.
- a protein antigen from *N.meningitidis* serogroup B, such as those in WO99/24578, WO99/36544, WO99/57280, WO00/22430, Tettelin *et al.* (2000) *Science* 287:1809-1815, Pizza *et al.* (2000) *Science* 287:1816-1820 and WO96/29412, with protein '287' and derivatives being particularly
- 20 preferred.
- an outer-membrane vesicle (OMV) preparation from *N.meningitidis* serogroup B, such as those disclosed in WO01/52885; Bjune *et al.* (1991) *Lancet* 338(8775):1093-1096; Fukasawa *et al.* (1999) *Vaccine* 17:2951-2958; Rosenqvist *et al.* (1998) *Dev. Biol. Stand.* 92:323-333 *etc.*
- a saccharide antigen from *N.meningitidis* serogroup A, C, W135 and/or Y, such as the
- 25 oligosaccharide disclosed in Costantino *et al.* (1992) *Vaccine* 10:691-698 from serogroup C [see also Costantino *et al.* (1999) *Vaccine* 17:1251-1263].
- a saccharide antigen from *Streptococcus pneumoniae* [e.g. Watson (2000) *Pediatr Infect Dis J* 19:331-332; Rubin (2000) *Pediatr Clin North Am* 47:269-285, v; Jedrzejewski (2001) *Microbiol Mol Biol Rev* 65:187-207].
- 30 – an antigen from hepatitis A virus, such as inactivated virus [e.g. Bell (2000) *Pediatr Infect Dis J* 19:1187-1188; Iwarson (1995) *APMIS* 103:321-326].
- an antigen from hepatitis B virus, such as the surface and/or core antigens [e.g. Gerlich *et al.* (1990) *Vaccine* 8 Suppl:S63-68 & 79-80].
- an antigen from hepatitis C virus [e.g. Hsu *et al.* (1999) *Clin Liver Dis* 3:901-915].
- 35 – an antigen from *Bordetella pertussis*, such as pertussis holotoxin (PT) and filamentous haemagglutinin (FHA) from *B.pertussis*, optionally also in combination with pertactin and/or

agglutinogens 2 and 3 [e.g. Gustafsson *et al.* (1996) *N. Engl. J. Med.* 334:349-355; Rappuoli *et al.* (1991) *TIBTECH* 9:232-238].

- a diphtheria antigen, such as a diphtheria toxoid [e.g. chapter 3 of *Vaccines* (1988) eds. Plotkin & Mortimer. ISBN 0-7216-1946-0] e.g. the CRM₁₉₇ mutant [e.g. Del Giudice *et al.* (1998) *Molecular Aspects of Medicine* 19:1-70].
- a tetanus antigen, such as a tetanus toxoid [e.g. chapter 4 of Plotkin & Mortimer].
- a saccharide antigen from *Haemophilus influenzae* B.
- an antigen from *N.gonorrhoeae* [e.g. WO99/24578, WO99/36544, WO99/57280].
- an antigen from *Chlamydia pneumoniae* [e.g. PCT/IB01/01445; Kalman *et al.* (1999) *Nature Genetics* 21:385-389; Read *et al.* (2000) *Nucleic Acids Res* 28:1397-406; Shirai *et al.* (2000) *J. Infect. Dis.* 181(Suppl 3):S524-S527; WO99/27105; WO00/27994; WO00/37494].
- an antigen from *Chlamydia trachomatis* [e.g. WO99/28475].
- an antigen from *Porphyromonas gingivalis* [e.g. Ross *et al.* (2001) *Vaccine* 19:4135-4142].
- polio antigen(s) [e.g. Sutter *et al.* (2000) *Pediatr Clin North Am* 47:287-308; Zimmerman & Spann (1999) *Am Fam Physician* 59:113-118, 125-126] such as IPV or OPV.
- rabies antigen(s) [e.g. Dreesen (1997) *Vaccine* 15 Suppl:S2-6] such as lyophilised inactivated virus [e.g. *MMWR Morb Mortal Wkly Rep* 1998 Jan 16;47(1):12, 19; RabAvertTM].
- measles, mumps and/or rubella antigens [e.g. chapters 9, 10 & 11 of Plotkin & Mortimer].
- influenza antigen(s) [e.g. chapter 19 of Plotkin & Mortimer], such as the haemagglutinin and/or neuraminidase surface proteins.
- an antigen from *Moraxella catarrhalis* [e.g. McMichael (2000) *Vaccine* 19 Suppl 1:S101-107].
- an antigen from *Staphylococcus aureus* [e.g. Kuroda *et al.* (2001) *Lancet* 357(9264):1225-1240; see also pages 1218-1219].

Where a saccharide or carbohydrate antigen is included, it is preferably conjugated to a carrier protein in order to enhance immunogenicity [e.g. Ramsay *et al.* (2001) *Lancet* 357(9251):195-196; Lindberg (1999) *Vaccine* 17 Suppl 2:S28-36; *Conjugate Vaccines* (eds. Cruse *et al.*) ISBN 3805549326, particularly vol. 10:48-114 *etc.*]. Preferred carrier proteins are bacterial toxins or toxoids, such as diphtheria or tetanus toxoids. The CRM₁₉₇ diphtheria toxoid is particularly preferred. Other suitable carrier proteins include the *N.meningitidis* outer membrane protein [e.g. EP-0372501], synthetic peptides [e.g. EP-0378881, EP-0427347], heat shock proteins [e.g. WO93/17712], pertussis proteins [e.g. WO98/58668; EP-0471177], protein D from *H.influenzae* [e.g. WO00/56360], toxin A or B from *C.difficile* [e.g. WO00/61761], *etc.* Any suitable conjugation reaction can be used, with any suitable linker where necessary.

Toxic protein antigens may be detoxified where necessary (e.g. detoxification of pertussis toxin by chemical and/or genetic means).

Where a diphtheria antigen is included in the composition it is preferred also to include tetanus antigen and pertussis antigens. Similarly, where a tetanus antigen is included it is preferred also to include diphtheria and pertussis antigens. Similarly, where a pertussis antigen is included it is preferred also to include diphtheria and tetanus antigens.

- 5 Antigens are preferably adsorbed to an aluminium salt.

Antigens in the composition will typically be present at a concentration of at least 1µg/ml each. In general, the concentration of any given antigen will be sufficient to elicit an immune response against that antigen.

The invention also provides compositions comprising two or more proteins of the present invention.

- 10 The two or more proteins may comprise GBS sequences or may comprise GAS and GBS sequences.

A summary of standard techniques and procedures which may be employed to perform the invention (e.g. to utilise the disclosed sequences for vaccination or diagnostic purposes) follows. This summary is not a limitation on the invention but, rather, gives examples that may be used, but are not required.

General

- 15 The practice of the present invention will employ, unless otherwise indicated, conventional techniques of molecular biology, microbiology, recombinant DNA, and immunology, which are within the skill of the art. Such techniques are explained fully in the literature eg. Sambrook *Molecular Cloning; A Laboratory Manual, Second Edition* (1989); *DNA Cloning, Volumes I and II* (D.N Glover ed. 1985); *Oligonucleotide Synthesis* (M.J. Gait ed, 1984); *Nucleic Acid Hybridization* (B.D. Hames & S.J. Higgins eds. 1984); *Transcription and Translation* (B.D. Hames & S.J. Higgins eds. 1984); *Animal Cell Culture* (R.I. Freshney ed. 1986); *Immobilized Cells and Enzymes* (IRL Press, 1986); B. Perbal, *A Practical Guide to Molecular Cloning* (1984); the *Methods in Enzymology* series (Academic Press, Inc.), especially volumes 154 & 155; *Gene Transfer Vectors for Mammalian Cells* (J.H. Miller and M.P. Calos eds. 1987, Cold Spring Harbor Laboratory); Mayer and Walker, eds. (1987), *Immunochemical*
- 20 *Methods in Cell and Molecular Biology* (Academic Press, London); Scopes, (1987) *Protein Purification: Principles and Practice*, Second Edition (Springer-Verlag, N.Y.), and *Handbook of Experimental Immunology, Volumes I-IV* (D.M. Weir and C. C. Blackwell eds 1986).

Standard abbreviations for nucleotides and amino acids are used in this specification.

Definitions

- 30 A composition containing X is "substantially free of" Y when at least 85% by weight of the total X+Y in the composition is X. Preferably, X comprises at least about 90% by weight of the total of X+Y in the composition, more preferably at least about 95% or even 99% by weight.
- The term "comprising" means "including" as well as "consisting" e.g. a composition "comprising" X may consist exclusively of X or may include something additional e.g. X + Y.
- 35 The term "heterologous" refers to two biological components that are not found together in nature. The components may be host cells, genes, or regulatory regions, such as promoters. Although the heterologous components are not found together in nature, they can function together, as when a promoter heterologous to a gene is operably linked to the gene. Another example is where a streptococcus sequence is heterologous to a mouse host cell. A further examples would be two epitopes from the same or different proteins which have been assembled in a single protein in an arrangement not found in nature

An "origin of replication" is a polynucleotide sequence that initiates and regulates replication of polynucleotides, such as an expression vector. The origin of replication behaves as an autonomous unit of polynucleotide replication within a cell, capable of replication under its own control. An origin of replication may be needed for a vector to replicate in a particular host cell. With certain origins of replication, an expression vector can be reproduced at a high copy number in the presence of the appropriate proteins within the cell. Examples of origins are the autonomously replicating sequences, which are effective in yeast; and the viral T-antigen, effective in COS-7 cells.

A "mutant" sequence is defined as DNA, RNA or amino acid sequence differing from but having sequence identity with the native or disclosed sequence. Depending on the particular sequence, the degree of sequence identity between the native or disclosed sequence and the mutant sequence is preferably greater than 50% (e.g. 60%, 70%, 80%, 90%, 95%, 99% or more, calculated using the Smith-Waterman algorithm as described above). As used herein, an "allelic variant" of a nucleic acid molecule, or region, for which nucleic acid sequence is provided herein is a nucleic acid molecule, or region, that occurs essentially at the same locus in the genome of another or second isolate, and that, due to natural variation caused by, for example, mutation or recombination, has a similar but not identical nucleic acid sequence. A coding region allelic variant typically encodes a protein having similar activity to that of the protein encoded by the gene to which it is being compared. An allelic variant can also comprise an alteration in the 5' or 3' untranslated regions of the gene, such as in regulatory control regions (e.g. see US patent 5,753,235).

Expression systems

The streptococcus nucleotide sequences can be expressed in a variety of different expression systems; for example those used with mammalian cells, baculoviruses, plants, bacteria, and yeast.

i. Mammalian Systems

Mammalian expression systems are known in the art. A mammalian promoter is any DNA sequence capable of binding mammalian RNA polymerase and initiating the downstream (3') transcription of a coding sequence (e.g. structural gene) into mRNA. A promoter will have a transcription initiating region, which is usually placed proximal to the 5' end of the coding sequence, and a TATA box, usually located 25-30 base pairs (bp) upstream of the transcription initiation site. The TATA box is thought to direct RNA polymerase II to begin RNA synthesis at the correct site. A mammalian promoter will also contain an upstream promoter element, usually located within 100 to 200 bp upstream of the TATA box. An upstream promoter element determines the rate at which transcription is initiated and can act in either orientation [Sambrook et al. (1989) "Expression of Cloned Genes in Mammalian Cells." In *Molecular Cloning: A Laboratory Manual*, 2nd ed.].

Mammalian viral genes are often highly expressed and have a broad host range; therefore sequences encoding mammalian viral genes provide particularly useful promoter sequences. Examples include the SV40 early promoter, mouse mammary tumor virus LTR promoter, adenovirus major late promoter (Ad MLP), and herpes simplex virus promoter. In addition, sequences derived from non-viral genes, such as the murine metallothionein gene, also provide useful promoter sequences. Expression may be either constitutive or regulated (inducible), depending on the promoter can be induced with glucocorticoid in hormone-responsive cells.

The presence of an enhancer element (enhancer), combined with the promoter elements described above, will usually increase expression levels. An enhancer is a regulatory DNA sequence that can stimulate transcription up to 1000-fold when linked to homologous or heterologous promoters, with synthesis beginning at the normal RNA start site. Enhancers are also active when they are placed upstream or downstream from the transcription initiation site, in either normal or flipped orientation, or at a distance of more than 1000 nucleotides from the promoter [Maniatis et al. (1987) *Science* 236:1237; Alberts et al. (1989) *Molecular Biology of the Cell*, 2nd ed.]. Enhancer elements derived from viruses may be particularly useful, because they usually have a broader host range. Examples include the SV40 early gene enhancer [Dijkema et al. (1985) *EMBO J.* 4:761] and the enhancer/promoters derived from the long terminal repeat (LTR) of the Rous Sarcoma Virus [Gorman et al. (1982b) *Proc. Natl. Acad. Sci.* 79:6777] and from human cytomegalovirus [Boshart et al. (1985) *Cell* 41:521]. Additionally, some enhancers are regulatable and become active only in the presence of an inducer, such as a hormone or metal ion [Sassone-Corsi and Borelli (1986) *Trends Genet.* 2:215; Maniatis et al. (1987) *Science* 236:1237].

A DNA molecule may be expressed intracellularly in mammalian cells. A promoter sequence may be directly linked with the DNA molecule, in which case the first amino acid at the N-terminus of the recombinant protein will always be a methionine, which is encoded by the ATG start codon. If desired, the N-terminus may be cleaved from the protein by *in vitro* incubation with cyanogen bromide.

Alternatively, foreign proteins can also be secreted from the cell into the growth media by creating chimeric DNA molecules that encode a fusion protein comprised of a leader sequence fragment that provides for secretion of the foreign protein in mammalian

cells. Preferably, there are processing sites encoded between the leader fragment and the foreign gene that can be cleaved either *in vivo* or *in vitro*. The leader sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the secretion of the protein from the cell. The adenovirus tripartite leader is an example of a leader sequence that provides for secretion of a foreign protein in mammalian cells.

- 5 Usually, transcription termination and polyadenylation sequences recognized by mammalian cells are regulatory regions located 3' to the translation stop codon and thus, together with the promoter elements, flank the coding sequence. The 3' terminus of the mature mRNA is formed by site-specific post-transcriptional cleavage and polyadenylation [Bimstiel et al. (1985) *Cell* 41:349; Proudfoot and Whitelaw (1988) "Termination and 3' end processing of eukaryotic RNA. In *Transcription and splicing* (ed. B.D. Hames and D.M. Glover); Proudfoot (1989) *Trends Biochem. Sci.* 14:105]. These sequences direct the transcription of an mRNA which can be translated into the polypeptide encoded by the DNA. Examples of transcription terminator/polyadenylation signals include those derived from SV40 [Sambrook et al. (1989) "Expression of cloned genes in cultured mammalian cells." In *Molecular Cloning: A Laboratory Manual*].

- Usually, the above described components, comprising a promoter, polyadenylation signal, and transcription termination sequence are put together into expression constructs. Enhancers, introns with functional splice donor and acceptor sites, and leader sequences may also be included in an expression construct, if desired. Expression constructs are often maintained in a replicon, such as an extrachromosomal element (eg. plasmids) capable of stable maintenance in a host, such as mammalian cells or bacteria. Mammalian replicon systems include those derived from animal viruses, which require trans-acting factors to replicate. For example, plasmids containing the replication systems of papovaviruses, such as SV40 [Gluzman (1981) *Cell* 23:175] or polyomavirus, replicate to extremely high copy number in the presence of the appropriate viral T antigen. Additional examples of mammalian replicons include those derived from bovine papillomavirus and Epstein-Barr virus. Additionally, the replicon may have two replication systems, thus allowing it to be maintained, for example, in mammalian cells for expression and in a prokaryotic host for cloning and amplification. Examples of such mammalian-bacteria shuttle vectors include pMT2 [Kaufman et al. (1989) *Mol. Cell. Biol.* 9:946] and pHEBO [Shimizu et al. (1986) *Mol. Cell. Biol.* 6:1074].

- 25 The transformation procedure used depends upon the host to be transformed. Methods for introduction of heterologous polynucleotides into mammalian cells are known in the art and include dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei.

- Mammalian cell lines available as hosts for expression are known in the art and include many immortalized cell lines available from the American Type Culture Collection (ATCC), including but not limited to, Chinese hamster ovary (CHO) cells, HeLa cells, baby hamster kidney (BHK) cells, monkey kidney cells (COS), human hepatocellular carcinoma cells (eg. Hep G2), and a number of other cell lines.

ii. Baculovirus Systems

- The polynucleotide encoding the protein can also be inserted into a suitable insect expression vector, and is operably linked to the control elements within that vector. Vector construction employs techniques which are known in the art. Generally, the components of the expression system include a transfer vector, usually a bacterial plasmid, which contains both a fragment of the baculovirus genome, and a convenient restriction site for insertion of the heterologous gene or genes to be expressed; a wild type baculovirus with a sequence homologous to the baculovirus-specific fragment in the transfer vector (this allows for the homologous recombination of the heterologous gene in to the baculovirus genome); and appropriate insect host cells and growth media.

- 40 After inserting the DNA sequence encoding the protein into the transfer vector, the vector and the wild type viral genome are transfected into an insect host cell where the vector and viral genome are allowed to recombine. The packaged recombinant virus is expressed and recombinant plaques are identified and purified. Materials and methods for baculovirus/insect cell expression systems are commercially available in kit form from, *inter alia*, Invitrogen, San Diego CA ("MaxBac" kit). These techniques are generally known to those skilled in the art and fully described in Summers and Smith, *Texas Agricultural Experiment Station Bulletin No. 1555* (1987) (hereinafter "Summers and Smith").

- Prior to inserting the DNA sequence encoding the protein into the baculovirus genome, the above described components, comprising a promoter, leader (if desired), coding sequence, and transcription termination sequence, are usually assembled into an intermediate transplacement construct (transfer vector). This may contain a single gene and operably linked regulatory elements; multiple genes, each with its own set of operably linked regulatory elements; or multiple genes, regulated by the same set of regulatory elements. Intermediate transplacement constructs are often maintained in a replicon, such as an extra-chromosomal

element (e.g. plasmids) capable of stable maintenance in a host, such as a bacterium. The replicon will have a replication system, thus allowing it to be maintained in a suitable host for cloning and amplification.

Currently, the most commonly used transfer vector for introducing foreign genes into AcNPV is pAc373. Many other vectors, known to those of skill in the art, have also been designed. These include, for example, pVL985 (which alters the polyhedrin start codon from ATG to ATT, and which introduces a BamHI cloning site 32 basepairs downstream from the ATT; see Luckow and Summers, *Virology* (1989) 17:31.

The plasmid usually also contains the polyhedrin polyadenylation signal (Miller et al. (1988) *Ann. Rev. Microbiol.*, 42:177) and a prokaryotic ampicillin-resistance (*amp*) gene and origin of replication for selection and propagation in *E.coli*.

Baculovirus transfer vectors usually contain a baculovirus promoter. A baculovirus promoter is any DNA sequence capable of binding a baculovirus RNA polymerase and initiating the downstream (5' to 3') transcription of a coding sequence (e.g. structural gene) into mRNA. A promoter will have a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation region usually includes an RNA polymerase binding site and a transcription initiation site. A baculovirus transfer vector may also have a second domain called an enhancer, which, if present, is usually distal to the structural gene. Expression may be either regulated or constitutive.

Structural genes, abundantly transcribed at late times in a viral infection cycle, provide particularly useful promoter sequences. Examples include sequences derived from the gene encoding the viral polyhedron protein, Friesen et al., (1986) "The Regulation of Baculovirus Gene Expression," in: *The Molecular Biology of Baculoviruses* (ed. Walter Doerfler); EPO Publ. Nos. 127 839 and 155 476; and the gene encoding the p10 protein, Vlak et al., (1988), *J. Gen. Virol.* 69:765.

DNA encoding suitable signal sequences can be derived from genes for secreted insect or baculovirus proteins, such as the baculovirus polyhedrin gene (Carbonell et al. (1988) *Gene*, 73:409). Alternatively, since the signals for mammalian cell posttranslational modifications (such as signal peptide cleavage, proteolytic cleavage, and phosphorylation) appear to be recognized by insect cells, and the signals required for secretion and nuclear accumulation also appear to be conserved between the invertebrate cells and vertebrate cells, leaders of non-insect origin, such as those derived from genes encoding human α -interferon, Maeda et al., (1985), *Nature* 315:592; human gastrin-releasing peptide, Lebacqz-Verheyden et al., (1988), *Molec. Cell. Biol.* 8:3129; human IL-2, Smith et al., (1985) *Proc. Natl Acad. Sci. USA*, 82:8404; mouse IL-3, (Miyajima et al., (1987) *Gene* 58:273; and human glucocerebrosidase, Martin et al. (1988) *DNA*, 7:99, can also be used to provide for secretion in insects.

A recombinant polypeptide or polypeptide may be expressed intracellularly or, if it is expressed with the proper regulatory sequences, it can be secreted. Good intracellular expression of nonfused foreign proteins usually requires heterologous genes that ideally have a short leader sequence containing suitable translation initiation signals preceding an ATG start signal. If desired, methionine at the N-terminus may be cleaved from the mature protein by *in vitro* incubation with cyanogen bromide.

Alternatively, recombinant polypeptides or proteins which are not naturally secreted can be secreted from the insect cell by creating chimeric DNA molecules that encode a fusion protein comprised of a leader sequence fragment that provides for secretion of the foreign protein in insects. The leader sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the translocation of the protein into the endoplasmic reticulum.

After insertion of the DNA sequence and/or the gene encoding the expression product precursor of the protein, an insect cell host is co-transformed with the heterologous DNA of the transfer vector and the genomic DNA of wild type baculovirus - usually by co-transfection. The promoter and transcription termination sequence of the construct will usually comprise a 2.5-kb section of the baculovirus genome. Methods for introducing heterologous DNA into the desired site in the baculovirus virus are known in the art. (See Summers and Smith *supra*; Ju et al. (1987); Smith et al., *Mol. Cell. Biol.* (1983) 3:2156; and Luckow and Summers (1989)). For example, the insertion can be into a gene such as the polyhedrin gene, by homologous double crossover recombination; insertion can also be into a restriction enzyme site engineered into the desired baculovirus gene. Miller et al., (1989), *Bioessays* 4:91. The DNA sequence, when cloned in place of the polyhedrin gene in the expression vector, is flanked both 5' and 3' by polyhedrin-specific sequences and is positioned downstream of the polyhedrin promoter.

The newly formed baculovirus expression vector is subsequently packaged into an infectious recombinant baculovirus. Homologous recombination occurs at low frequency (between about 1% and about 5%); thus, the majority of the virus produced after cotransfection is still wild-type virus. Therefore, a method is necessary to identify recombinant viruses. An advantage of the expression system is a visual screen allowing recombinant viruses to be distinguished. The polyhedrin protein, which is produced by the native virus, is produced at very high levels in the nuclei of infected cells at late times after viral infection. Accumulated polyhedrin protein forms occlusion bodies that also contain embedded particles. These occlusion bodies, up to 15 μ m in size, are

highly refractile, giving them a bright shiny appearance that is readily visualized under the light microscope. Cells infected with recombinant viruses lack occlusion bodies. To distinguish recombinant virus from wild-type virus, the transfection supernatant is plaqued onto a monolayer of insect cells by techniques known to those skilled in the art. Namely, the plaques are screened under the light microscope for the presence (indicative of wild-type virus) or absence (indicative of recombinant virus) of occlusion bodies. "Current Protocols in Microbiology" Vol. 2 (Ausubel et al. eds) at 16.8 (Supp. 10, 1990); Summers and Smith, *supra*; Miller et al. (1989).

Recombinant baculovirus expression vectors have been developed for infection into several insect cells. For example, recombinant baculoviruses have been developed for, *inter alia*: *Aedes aegypti*, *Autographa californica*, *Bombyx mori*, *Drosophila melanogaster*, *Spodoptera frugiperda*, and *Trichoplusia ni* (WO 89/046699; Carbonell et al., (1985) *J. Virol.* 56:153; Wright (1986) *Nature* 321:718; Smith et al., (1983) *Mol. Cell. Biol.* 3:2156; and see generally, Fraser, et al. (1989) *In Vitro Cell. Dev. Biol.* 25:225).

Cells and cell culture media are commercially available for both direct and fusion expression of heterologous polypeptides in a baculovirus/expression system; cell culture technology is generally known to those skilled in the art. See, eg. Summers and Smith *supra*.

The modified insect cells may then be grown in an appropriate nutrient medium, which allows for stable maintenance of the plasmid(s) present in the modified insect host. Where the expression product gene is under inducible control, the host may be grown to high density, and expression induced. Alternatively, where expression is constitutive, the product will be continuously expressed into the medium and the nutrient medium must be continuously circulated, while removing the product of interest and augmenting depleted nutrients. The product may be purified by such techniques as chromatography, eg. HPLC, affinity chromatography, ion exchange chromatography, etc.; electrophoresis; density gradient centrifugation; solvent extraction, etc. As appropriate, the product may be further purified, as required, so as to remove substantially any insect proteins which are also present in the medium, so as to provide a product which is at least substantially free of host debris, eg. proteins, lipids and polysaccharides.

In order to obtain protein expression, recombinant host cells derived from the transformants are incubated under conditions which allow expression of the recombinant protein encoding sequence. These conditions will vary, dependent upon the host cell selected. However, the conditions are readily ascertainable to those of ordinary skill in the art, based upon what is known in the art.

iii. Plant Systems

There are many plant cell culture and whole plant genetic expression systems known in the art. Exemplary plant cellular genetic expression systems include those described in patents, such as: US 5,693,506; US 5,659,122; and US 5,608,143. Additional examples of genetic expression in plant cell culture has been described by Zenk, *Phytochemistry* 30:3861-3863 (1991). Descriptions of plant protein signal peptides may be found in addition to the references described above in Vaulcombe et al., *Mol. Gen. Genet.* 209:33-40 (1987); Chandler et al., *Plant Molecular Biology* 3:407-418 (1984); Rogers, *J. Biol. Chem.* 260:3731-3738 (1985); Rothstein et al., *Gene* 55:353-356 (1987); Whittier et al., *Nucleic Acids Research* 15:2515-2535 (1987); Wirsal et al., *Molecular Microbiology* 3:3-14 (1989); Yu et al., *Gene* 122:247-253 (1992). A description of the regulation of plant gene expression by the phytohormone, gibberellic acid and secreted enzymes induced by gibberellic acid can be found in R.L. Jones and J. MacMillan, Gibberellins: in: *Advanced Plant Physiology*, Malcolm B. Wilkins, ed., 1984 Pitman Publishing Limited, London, pp. 21-52. References that describe other metabolically-regulated genes: Sheen, *Plant Cell*, 2:1027-1038(1990); Maas et al., *EMBO J.* 9:3447-3452 (1990); Benkel and Hickey, *Proc. Natl. Acad. Sci.* 84:1337-1339 (1987).

Typically, using techniques known in the art, a desired polynucleotide sequence is inserted into an expression cassette comprising genetic regulatory elements designed for operation in plants. The expression cassette is inserted into a desired expression vector with companion sequences upstream and downstream from the expression cassette suitable for expression in a plant host. The companion sequences will be of plasmid or viral origin and provide necessary characteristics to the vector to permit the vectors to move DNA from an original cloning host, such as bacteria, to the desired plant host. The basic bacterial/plant vector construct will preferably provide a broad host range prokaryote replication origin; a prokaryote selectable marker; and, for Agrobacterium transformations, T DNA sequences for Agrobacterium-mediated transfer to plant chromosomes. Where the heterologous gene is not readily amenable to detection, the construct will preferably also have a selectable marker gene suitable for determining if a plant cell has been transformed. A general review of suitable markers, for example for the members of the grass family, is found in Wilmink and Dons, 1993, *Plant Mol. Biol. Repts*, 11(2):165-185.

Sequences suitable for permitting integration of the heterologous sequence into the plant genome are also recommended. These might include transposon sequences and the like for homologous recombination as well as 'IT' sequences which permit random insertion of a heterologous expression cassette into a plant genome. Suitable prokaryote selectable markers include resistance toward antibiotics such as ampicillin or tetracycline. Other DNA sequences encoding additional functions may also be present in the vector, as is known in the art.

The nucleic acid molecules of the subject invention may be included into an expression cassette for expression of the protein(s) of interest. Usually, there will be only one expression cassette, although two or more are feasible. The recombinant expression cassette will contain in addition to the heterologous protein encoding sequence the following elements, a promoter region, plant 5' untranslated sequences, initiation codon depending upon whether or not the structural gene comes equipped with one, and a transcription and translation termination sequence. Unique restriction enzyme sites at the 5' and 3' ends of the cassette allow for easy insertion into a pre-existing vector.

A heterologous coding sequence may be for any protein relating to the present invention. The sequence encoding the protein of interest will encode a signal peptide which allows processing and translocation of the protein, as appropriate, and will usually lack any sequence which might result in the binding of the desired protein of the invention to a membrane. Since, for the most part, the transcriptional initiation region will be for a gene which is expressed and translocated during germination, by employing the signal peptide which provides for translocation, one may also provide for translocation of the protein of interest. In this way, the protein(s) of interest will be translocated from the cells in which they are expressed and may be efficiently harvested. Typically secretion in seeds are across the aleurone or scutellar epithelium layer into the endosperm of the seed. While it is not required that the protein be secreted from the cells in which the protein is produced, this facilitates the isolation and purification of the recombinant protein.

Since the ultimate expression of the desired gene product will be in a eucaryotic cell it is desirable to determine whether any portion of the cloned gene contains sequences which will be processed out as introns by the host's spliceosome machinery. If so, site-directed mutagenesis of the "intron" region may be conducted to prevent losing a portion of the genetic message as a false intron code, Reed and Maniatis, *Cell* 41:95-105, 1985.

The vector can be microinjected directly into plant cells by use of micropipettes to mechanically transfer the recombinant DNA. Crossway, *Mol. Gen. Genet.*, 202:179-185, 1985. The genetic material may also be transferred into the plant cell by using polyethylene glycol, Krens, et al., *Nature*, 296, 72-74, 1982. Another method of introduction of nucleic acid segments is high velocity ballistic penetration by small particles with the nucleic acid either within the matrix of small beads or particles, or on the surface, Klein, et al., *Nature*, 327, 70-73, 1987 and Knudsen and Muller, 1991, *Planta*, 185:330-336 teaching particle bombardment of barley endosperm to create transgenic barley. Yet another method of introduction would be fusion of protoplasts with other entities, either minicells, cells, lysosomes or other fusible lipid-surfaced bodies, Fraley, et al., *Proc. Natl. Acad. Sci. USA*, 79, 1859-1863, 1982.

The vector may also be introduced into the plant cells by electroporation. (Fromm et al., *Proc. Natl. Acad. Sci. USA* 82:5824, 1985). In this technique, plant protoplasts are electroporated in the presence of plasmids containing the gene construct. Electrical impulses of high field strength reversibly permeabilize biomembranes allowing the introduction of the plasmids. Electroporated plant protoplasts reform the cell wall, divide, and form plant callus.

All plants from which protoplasts can be isolated and cultured to give whole regenerated plants can be transformed by the present invention so that whole plants are recovered which contain the transferred gene. It is known that practically all plants can be regenerated from cultured cells or tissues, including but not limited to all major species of sugarcane, sugar beet, cotton, fruit and other trees, legumes and vegetables. Some suitable plants include, for example, species from the genera *Fragaria*, *Lotus*, *Medicago*, *Onobrychis*, *Trifolium*, *Trigonella*, *Vigna*, *Citrus*, *Linum*, *Geranium*, *Manihot*, *Daucus*, *Arabidopsis*, *Brassica*, *Raphanus*, *Sinapis*, *Atropa*, *Capsicum*, *Datura*, *Hyoscyamus*, *Lycopersion*, *Nicotiana*, *Solanum*, *Petunia*, *Digitalis*, *Majorana*, *Cichorium*, *Helianthus*, *Lactuca*, *Bromus*, *Asparagus*, *Antirrhinum*, *Hererocallis*, *Nemesia*, *Pelargonium*, *Panicum*, *Pennisetum*, *Ranunculus*, *Senecio*, *Salpiglossis*, *Cucumis*, *Browaalia*, *Glycine*, *Lolium*, *Zea*, *Triticum*, *Sorghum*, and *Datura*.

Means for regeneration vary from species to species of plants, but generally a suspension of transformed protoplasts containing copies of the heterologous gene is first provided. Callus tissue is formed and shoots may be induced from callus and subsequently rooted. Alternatively, embryo formation can be induced from the protoplast suspension. These embryos germinate as natural embryos to form plants. The culture media will generally contain various amino acids and hormones, such as auxin and cytokinins. It is also advantageous to add glutamic acid and proline to the medium, especially for such species as corn and alfalfa. Shoots and

roots normally develop simultaneously. Efficient regeneration will depend on the medium, on the genotype, and on the history of the culture. If these three variables are controlled, then regeneration is fully reproducible and repeatable.

In some plant cell culture systems, the desired protein of the invention may be excreted or alternatively, the protein may be extracted from the whole plant. Where the desired protein of the invention is secreted into the medium, it may be collected. Alternatively, the embryos and embryoless-half seeds or other plant tissue may be mechanically disrupted to release any secreted protein between cells and tissues. The mixture may be suspended in a buffer solution to retrieve soluble proteins. Conventional protein isolation and purification methods will be then used to purify the recombinant protein. Parameters of time, temperature pH, oxygen, and volumes will be adjusted through routine methods to optimize expression and recovery of heterologous protein.

iv. Bacterial Systems

- 10 Bacterial expression techniques are known in the art. A bacterial promoter is any DNA sequence capable of binding bacterial RNA polymerase and initiating the downstream (3') transcription of a coding sequence (eg. structural gene) into mRNA. A promoter will have a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation region usually includes an RNA polymerase binding site and a transcription initiation site. A bacterial promoter may also have a second domain called an operator, that may overlap an adjacent RNA polymerase binding site at which RNA synthesis begins. The operator permits negative regulated (inducible) transcription, as a gene repressor protein may bind the operator and thereby inhibit transcription of a specific gene. Constitutive expression may occur in the absence of negative regulatory elements, such as the operator. In addition, positive regulation may be achieved by a gene activator protein binding sequence, which, if present is usually proximal (5') to the RNA polymerase binding sequence. An example of a gene activator protein is the catabolite activator protein (CAP), which helps initiate transcription of the lac operon in *Escherichia coli* (*E. coli*) [Raibaud *et al.* (1984) *Annu. Rev. Genet.* 18:173]. Regulated expression may therefore be either positive or negative, thereby either enhancing or reducing transcription.

Sequences encoding metabolic pathway enzymes provide particularly useful promoter sequences. Examples include promoter sequences derived from sugar metabolizing enzymes, such as galactose, lactose (*lac*) [Chang *et al.* (1977) *Nature* 198:1056], and maltose. Additional examples include promoter sequences derived from biosynthetic enzymes such as tryptophan (*trp*) [Goeddel *et al.* (1980) *Nuc. Acids Res.* 8:4057; Yelverton *et al.* (1981) *Nucl. Acids Res.* 9:731; US patent 4,738,921; EP-A-0036776 and EP-A-0121775]. The *g-lotamase* (*bla*) promoter system [Weissmann (1981) "The cloning of interferon and other mistakes." In *Interferon* 3 (ed. I. Gresser)], bacteriophage lambda PL [Shimatake *et al.* (1981) *Nature* 292:128] and T5 [US patent 4,689,406] promoter systems also provide useful promoter sequences.

- 25 In addition, synthetic promoters which do not occur in nature also function as bacterial promoters. For example, transcription activation sequences of one bacterial or bacteriophage promoter may be joined with the operon sequences of another bacterial or bacteriophage promoter, creating a synthetic hybrid promoter [US patent 4,551,433]. For example, the *tac* promoter is a hybrid *trp-lac* promoter comprised of both *trp* promoter and *lac* operon sequences that is regulated by the *lac* repressor [Amann *et al.* (1983) *Gene* 25:167; de Boer *et al.* (1983) *Proc. Natl. Acad. Sci.* 80:21]. Furthermore, a bacterial promoter can include naturally occurring promoters of non-bacterial origin that have the ability to bind bacterial RNA polymerase and initiate transcription. A naturally occurring promoter of non-bacterial origin can also be coupled with a compatible RNA polymerase to produce high levels of expression of some genes in prokaryotes. The bacteriophage T7 RNA polymerase/promoter system is an example of a coupled promoter system [Studier *et al.* (1986) *J. Mol. Biol.* 189:113; Tabor *et al.* (1985) *Proc Natl. Acad. Sci.* 82:1074]. In addition, a hybrid promoter can also be comprised of a bacteriophage promoter and an *E.coli* operator region (EPO-A-0 267 851).
- 40 In addition to a functioning promoter sequence, an efficient ribosome binding site is also useful for the expression of foreign genes in prokaryotes. In *E.coli*, the ribosome binding site is called the Shine-Dalgarno (SD) sequence and includes an initiation codon (ATG) and a sequence 3-9 nucleotides in length located 3-11 nucleotides upstream of the initiation codon [Shine *et al.* (1975) *Nature* 254:34]. The SD sequence is thought to promote binding of mRNA to the ribosome by the pairing of bases between the SD sequence and the 3' end of *E.coli* 16S rRNA [Steitz *et al.* (1979) "Genetic signals and nucleotide sequences in messenger RNA." In *Biological Regulation and Development: Gene Expression* (ed. R.F. Goldberg)]. To express eukaryotic genes and prokaryotic genes with weak ribosome-binding site [Sambrook *et al.* (1989) "Expression of cloned genes in *Escherichia coli*." In *Molecular Cloning: A Laboratory Manual*].

A DNA molecule may be expressed intracellularly. A promoter sequence may be directly linked with the DNA molecule, in which case the first amino acid at the N-terminus will always be a methionine, which is encoded by the ATG start codon. If desired, methionine at the N-terminus may be cleaved from the protein by *in vitro* incubation with cyanogen bromide or by either *in vivo* or *in vitro* incubation with a bacterial methionine N-terminal peptidase (EP-A-0 219 237).

Fusion proteins provide an alternative to direct expression. Usually, a DNA sequence encoding the N-terminal portion of an endogenous bacterial protein, or other stable protein, is fused to the 5' end of heterologous coding sequences. Upon expression, this construct will provide a fusion of the two amino acid sequences. For example, the bacteriophage lambda cell gene can be linked at the 5' terminus of a foreign gene and expressed in bacteria. The resulting fusion protein preferably retains a site for a processing enzyme (factor Xa) to cleave the bacteriophage protein from the foreign gene [Nagai *et al.* (1984) *Nature* 309:810]. Fusion proteins can also be made with sequences from the *lacZ* [Jia *et al.* (1987) *Gene* 60:197], *trpE* [Allen *et al.* (1987) *J. Biotechnol.* 5:93; Makoff *et al.* (1989) *J. Gen. Microbiol.* 135:11], and *Chey* [EP-A-0 324 647] genes. The DNA sequence at the junction of the two amino acid sequences may or may not encode a cleavable site. Another example is a ubiquitin fusion protein. Such a fusion protein is made with the ubiquitin region that preferably retains a site for a processing enzyme (e.g. ubiquitin specific processing-protease) to cleave the ubiquitin from the foreign protein. Through this method, native foreign protein can be isolated [Miller *et al.* (1989) *BioTechnology* 7:698].

Alternatively, foreign proteins can also be secreted from the cell by creating chimeric DNA molecules that encode a fusion protein comprised of a signal peptide sequence fragment that provides for secretion of the foreign protein in bacteria [US patent 4,336,336]. The signal sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the secretion of the protein from the cell. The protein is either secreted into the growth media (gram-positive bacteria) or into the periplasmic space, located between the inner and outer membrane of the cell (gram-negative bacteria). Preferably there are processing sites, which can be cleaved either *in vivo* or *in vitro* encoded between the signal peptide fragment and the foreign gene.

DNA encoding suitable signal sequences can be derived from genes for secreted bacterial proteins, such as the *E. coli* outer membrane protein gene (*ompA*) [Masui *et al.* (1983), in: *Experimental Manipulation of Gene Expression*; Ghayeb *et al.* (1984) *EMBO J.* 3:2437] and the *E. coli* alkaline phosphatase signal sequence (*phoA*) [Oka *et al.* (1985) *Proc. Natl. Acad. Sci.* 82:7212]. As an additional example, the signal sequence of the alpha-amylase gene from various *Bacillus* strains can be used to secrete heterologous proteins from *B. subtilis* [Palva *et al.* (1982) *Proc. Natl. Acad. Sci. USA* 79:5582; EP-A-0 244 042].

Usually, transcription termination sequences recognized by bacteria are regulatory regions located 3' to the translation stop codon, and thus together with the promoter flank the coding sequence. These sequences direct the transcription of an mRNA which can be translated into the polypeptide encoded by the DNA. Transcription termination sequences frequently include DNA sequences of about 50 nucleotides capable of forming stem loop structures that aid in terminating transcription. Examples include transcription termination sequences derived from genes with strong promoters, such as the *trp* gene in *E. coli* as well as other biosynthetic genes.

Usually, the above described components, comprising a promoter, signal sequence (if desired), coding sequence of interest, and transcription termination sequence, are put together into expression constructs. Expression constructs are often maintained in a replicon, such as an extrachromosomal element (e.g. plasmids) capable of stable maintenance in a host, such as bacteria. The replicon will have a replication system, thus allowing it to be maintained in a prokaryotic host either for expression or for cloning and amplification. In addition, a replicon may be either a high or low copy number plasmid. A high copy number plasmid will generally have a copy number ranging from about 5 to about 200, and usually about 10 to about 150. A host containing a high copy number plasmid will preferably contain at least about 10, and more preferably at least about 20 plasmids. Either a high or low copy number vector may be selected, depending upon the effect of the vector and the foreign protein on the host.

Alternatively, the expression constructs can be integrated into the bacterial genome with an integrating vector. Integrating vectors usually contain at least one sequence homologous to the bacterial chromosome that allows the vector to integrate. Integrations appear to result from recombinations between homologous DNA in the vector and the bacterial chromosome. For example, integrating vectors constructed with DNA from various *Bacillus* strains integrate into the *Bacillus* chromosome (EP-A-0 127 328). Integrating vectors may also be comprised of bacteriophage or transposon sequences.

Usually, extrachromosomal and integrating expression constructs may contain selectable markers to allow for the selection of bacterial strains that have been transformed. Selectable markers can be expressed in the bacterial host and may include genes which render bacteria resistant to drugs such as ampicillin, chloramphenicol, erythromycin, kanamycin (neomycin), and tetracycline [Davies *et al.* (1978) *Annu. Rev. Microbiol.* 32:469]. Selectable markers may also include biosynthetic genes, such as those in the histidine, tryptophan, and leucine biosynthetic pathways.

Alternatively, some of the above described components can be put together in transformation vectors. Transformation vectors are usually comprised of a selectable marker that is either maintained in a replicon or developed into an integrating vector, as described above.

Expression and transformation vectors, either extra-chromosomal replicons or integrating vectors, have been developed for transformation into many bacteria. For example, expression vectors have been developed for, *inter alia*, the following bacteria: *Bacillus subtilis* [Palva *et al.* (1982) *Proc. Natl. Acad. Sci. USA* 79:5582; EP-A-0 036 259 and EP-A-0 063 953; WO 84/04541]; *Escherichia coli* [Shimatake *et al.* (1981) *Nature* 292:128; Amann *et al.* (1985) *Gene* 40:183; Studier *et al.* (1986) *J. Mol. Biol.* 189:113; EP-A-0 036 776, EP-A-0 136 829 and EP-A-0 136 907]; *Streptococcus cremoris* [Powell *et al.* (1988) *Appl. Environ. Microbiol.* 54:655]; *Streptococcus lividans* [Powell *et al.* (1988) *Appl. Environ. Microbiol.* 54:655]; *Streptomyces lividans* [US patent 4,745,056].

Methods of introducing exogenous DNA into bacterial hosts are well-known in the art, and usually include either the transformation of bacteria treated with CaCl_2 or other agents, such as divalent cations and DMSO. DNA can also be introduced into bacterial cells by electroporation. Transformation procedures usually vary with the bacterial species to be transformed. See eg. [Masson *et al.* (1989) *FEMS Microbiol. Lett.* 60:273; Palva *et al.* (1982) *Proc. Natl. Acad. Sci. USA* 79:5582; EP-A-0 036 259 and EP-A-0 063 953; WO 84/04541, *Bacillus*], [Miller *et al.* (1988) *Proc. Natl. Acad. Sci.* 85:856; Wang *et al.* (1990) *J. Bacteriol.* 172:949, *Campylobacter*], [Cohen *et al.* (1973) *Proc. Natl. Acad. Sci.* 69:2110; Dower *et al.* (1988) *Nucleic Acids Res.* 16:6127; Kushner (1978) "An improved method for transformation of *Escherichia coli* with ColE1-derived plasmids. In *Genetic Engineering: Proceedings of the International Symposium on Genetic Engineering* (eds. H.W. Boyer and S. Nicosia); Mandel *et al.* (1970) *J. Mol. Biol.* 53:159; Taketo (1988) *Biochim. Biophys. Acta* 949:318; *Escherichia*], [Chassy *et al.* (1987) *FEMS Microbiol. Lett.* 44:173 *Lactobacillus*]; [Fiedler *et al.* (1988) *Anal. Biochem.* 170:38, *Pseudomonas*]; [Augustin *et al.* (1990) *FEMS Microbiol. Lett.* 66:203, *Staphylococcus*], [Barany *et al.* (1980) *J. Bacteriol.* 144:698; Harlander (1987) "Transformation of *Streptococcus lactis* by electroporation, in: *Streptococcal Genetics* (ed. J. Ferretti and R. Curtiss III); Perry *et al.* (1981) *Infect. Immun.* 32:1295; Powell *et al.* (1988) *Appl. Environ. Microbiol.* 54:655; Somkuti *et al.* (1987) *Proc. 4th Eur. Cong. Biotechnology* 1:412, *Streptococcus*].

v. Yeast Expression

Yeast expression systems are also known to one of ordinary skill in the art. A yeast promoter is any DNA sequence capable of binding yeast RNA polymerase and initiating the downstream (3') transcription of a coding sequence (eg. structural gene) into mRNA. A promoter will have a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation region usually includes an RNA polymerase binding site (the "TATA Box") and a transcription initiation site. A yeast promoter may also have a second domain called an upstream activator sequence (UAS), which, if present, is usually distal to the structural gene. The UAS permits regulated (inducible) expression. Constitutive expression occurs in the absence of a UAS. Regulated expression may be either positive or negative, thereby either enhancing or reducing transcription.

Yeast is a fermenting organism with an active metabolic pathway, therefore sequences encoding enzymes in the metabolic pathway provide particularly useful promoter sequences. Examples include alcohol dehydrogenase (ADH) (EP-A-0 284 044), endolase, glucokinase, glucose-6-phosphate isomerase, glyceraldehyde-3-phosphate-dehydrogenase (GAP or GAPDH), hexokinase, phosphofructokinase, 3-phosphoglycerate mutase, and pyruvate kinase (PyK) (EPO-A-0 329 203). The yeast *PHO5* gene, encoding acid phosphatase, also provides useful promoter sequences [Myranohara *et al.* (1983) *Proc. Natl. Acad. Sci. USA* 80:1].

In addition, synthetic promoters which do not occur in nature also function as yeast promoters. For example, UAS sequences of one yeast promoter may be joined with the transcription activation region of another yeast promoter, creating a synthetic hybrid promoter. Examples of such hybrid promoters include the ADH regulatory sequence linked to the GAP transcription activation region (US Patent Nos. 4,876,197 and 4,880,734). Other examples of hybrid promoters include promoters which consist of the regulatory sequences of either the *ADH2*, *GAL4*, *GAL10*, OR *PHO5* genes, combined with the transcriptional activation region of a glycolytic enzyme gene such as GAP or PyK (EP-A-0 164 556). Furthermore, a yeast promoter can include naturally occurring promoters of non-yeast origin that have the ability to bind yeast RNA polymerase and initiate transcription. Examples of such promoters include, *inter alia*, [Cohen *et al.* (1980) *Proc. Natl. Acad. Sci. USA* 77:1078; Henikoff *et al.* (1981) *Nature* 283:835; Hollenberg *et al.* (1981) *Curr. Topics Microbiol. Immunol.* 96:119; Hollenberg *et al.* (1979) "The Expression of Bacterial Antibiotic Resistance Genes in the Yeast *Saccharomyces cerevisiae*," in: *Plasmids of Medical, Environmental and Commercial Importance* (eds. K.N. Timmis and A. Puhler); Mercerau-Puigalon *et al.* (1980) *Gene* 11:163; Panthier *et al.* (1980) *Curr. Genet.* 2:109].

A DNA molecule may be expressed intracellularly in yeast. A promoter sequence may be directly linked with the DNA molecule, in which case the first amino acid at the N-terminus of the recombinant protein will always be a methionine, which is encoded by

the ATG start codon. If desired, methionine at the N-terminus may be cleaved from the protein by *in vitro* incubation with cyanogen bromide.

Fusion proteins provide an alternative for yeast expression systems, as well as in mammalian, baculovirus, and bacterial expression systems. Usually, a DNA sequence encoding the N-terminal portion of an endogenous yeast protein, or other stable protein, is fused to the 5' end of heterologous coding sequences. Upon expression, this construct will provide a fusion of the two amino acid sequences. For example, the yeast or human superoxide dismutase (SOD) gene, can be linked at the 5' terminus of a foreign gene and expressed in yeast. The DNA sequence at the junction of the two amino acid sequences may or may not encode a cleavable site. See eg. EP-A-0 196 056. Another example is a ubiquitin fusion protein. Such a fusion protein is made with the ubiquitin region that preferably retains a site for a processing enzyme (eg. ubiquitin-specific processing protease) to cleave the ubiquitin from the foreign protein. Through this method, therefore, native foreign protein can be isolated (eg. WO88/024066).

Alternatively, foreign proteins can also be secreted from the cell into the growth media by creating chimeric DNA molecules that encode a fusion protein comprised of a leader sequence fragment that provide for secretion in yeast of the foreign protein. Preferably, there are processing sites encoded between the leader fragment and the foreign gene that can be cleaved either *in vivo* or *in vitro*. The leader sequence fragment usually encodes a signal peptide comprised of hydrophobic amino acids which direct the secretion of the protein from the cell.

DNA encoding suitable signal sequences can be derived from genes for secreted yeast proteins, such as the yeast invertase gene (EP-A-0 012 873; JPO. 62,096,086) and the A-factor gene (US patent 4,588,684). Alternatively, leaders of non-yeast origin, such as an interferon leader, exist that also provide for secretion in yeast (EP-A-0 060 057).

A preferred class of secretion leaders are those that employ a fragment of the yeast alpha-factor gene, which contains both a "pre" signal sequence, and a "pro" region. The types of alpha-factor fragments that can be employed include the full-length pre-pro alpha factor leader (about 83 amino acid residues) as well as truncated alpha-factor leaders (usually about 25 to about 50 amino acid residues) (US Patents 4,546,083 and 4,870,008; EP-A-0 324 274). Additional leaders employing an alpha-factor leader fragment that provides for secretion include hybrid alpha-factor leaders made with a presequence of a first yeast, but a pro-region from a second yeast alphafactor. (eg. see WO 89/02463.)

Usually, transcription termination sequences recognized by yeast are regulatory regions located 3' to the translation stop codon, and thus together with the promoter flank the coding sequence. These sequences direct the transcription of an mRNA which can be translated into the polypeptide encoded by the DNA. Examples of transcription terminator sequence and other yeast-recognized termination sequences, such as those coding for glycolytic enzymes.

Usually, the above described components, comprising a promoter, leader (if desired), coding sequence of interest, and transcription termination sequence, are put together into expression constructs. Expression constructs are often maintained in a replicon, such as an extrachromosomal element (eg. plasmids) capable of stable maintenance in a host, such as yeast or bacteria. The replicon may have two replication systems, thus allowing it to be maintained, for example, in yeast for expression and in a prokaryotic host for cloning and amplification. Examples of such yeast-bacteria shuttle vectors include YEp24 [Botstein *et al.* (1979) *Gene* 8:17-24], pCI/1 [Brake *et al.* (1984) *Proc. Natl. Acad. Sci USA* 81:4642-4646], and YRp17 [Stinchcomb *et al.* (1982) *J. Mol. Biol.* 158:157]. In addition, a replicon may be either a high or low copy number plasmid. A high copy number plasmid will generally have a copy number ranging from about 5 to about 200, and usually about 10 to about 150. A host containing a high copy number plasmid will preferably have at least about 10, and more preferably at least about 20. Enter a high or low copy number vector may be selected, depending upon the effect of the vector and the foreign protein on the host. See eg. Brake *et al.*, *supra*.

Alternatively, the expression constructs can be integrated into the yeast genome with an integrating vector. Integrating vectors usually contain at least one sequence homologous to a yeast chromosome that allows the vector to integrate, and preferably contain two homologous sequences flanking the expression construct. Integrations appear to result from recombinations between homologous DNA in the vector and the yeast chromosome [Orr-Weaver *et al.* (1983) *Methods in Enzymol.* 101:228-245]. An integrating vector may be directed to a specific locus in yeast by selecting the appropriate homologous sequence for inclusion in the vector. See Orr-Weaver *et al.*, *supra*. One or more expression construct may integrate, possibly affecting levels of recombinant protein produced [Rine *et al.* (1983) *Proc. Natl. Acad. Sci. USA* 80:6750]. The chromosomal sequences included in the vector can occur either as a single segment in the vector, which results in the integration of the entire vector, or two segments homologous to adjacent segments in the chromosome and flanking the expression construct in the vector, which can result in the stable integration of only the expression construct.

Usually, extrachromosomal and integrating expression constructs may contain selectable markers to allow for the selection of yeast strains that have been transformed. Selectable markers may include biosynthetic genes that can be expressed in the yeast host, such as *ADE2*, *HIS4*, *LEU2*, *TRP1*, and *ALG7*, and the G418 resistance gene, which confer resistance in yeast cells to tunicamycin and G418, respectively. In addition, a suitable selectable marker may also provide yeast with the ability to grow in the presence of toxic compounds, such as metal. For example, the presence of *CUP1* allows yeast to grow in the presence of copper ions [Butt *et al.* (1987) *Microbiol. Rev.* 51:351].

Alternatively, some of the above described components can be put together into transformation vectors. Transformation vectors are usually comprised of a selectable marker that is either maintained in a replicon or developed into an integrating vector, as described above.

- 10 Expression and transformation vectors, either extrachromosomal replicons or integrating vectors, have been developed for transformation into many yeasts. For example, expression vectors have been developed for, *inter alia*, the following yeasts: *Candida albicans* [Kurtz, *et al.* (1986) *Mol. Cell. Biol.* 6:142], *Candida maltosa* [Kunze, *et al.* (1985) *J. Basic Microbiol.* 25:141], *Hansenula polymorpha* [Gleeson, *et al.* (1986) *J. Gen. Microbiol.* 132:3459; Roggenkamp *et al.* (1986) *Mol. Gen. Genet.* 202:302], *Kluyveromyces fragilis* [Das, *et al.* (1984) *J. Bacteriol.* 158:1165], *Kluyveromyces fragilis* [De Louvencourt *et al.* (1983) *J. Bacteriol.* 154:737; Van den Berg *et al.* (1990) *BioTechnology* 8:135], *Pichia guilliermondii* [Kunze *et al.* (1985) *J. Basic Microbiol.* 25:141], *Pichia pastoris* [Cregg, *et al.* (1985) *Mol. Cell. Biol.* 5:3376; US Patent Nos. 4,837,148 and 4,929,555], *Saccharomyces cerevisiae* [Hinnen *et al.* (1978) *Proc. Natl. Acad. Sci. USA* 75:1929; Ito *et al.* (1983) *J. Bacteriol.* 153:163], *Schizosaccharomyces pombe* [Beach and Nurse (1981) *Nature* 300:706], and *Yarrowia lipolytica* [Davidow, *et al.* (1985) *Curr. Genet.* 10:380471 Gaillardin, *et al.* (1985) *Curr. Genet.* 10:49].
- 20 Methods of introducing exogenous DNA into yeast hosts are well-known in the art, and usually include either the transformation of spheroplasts or of intact yeast cells treated with alkali cations. Transformation procedures usually vary with the yeast species to be transformed. See eg. [Kurtz, *et al.* (1986) *Mol. Cell. Biol.* 6:142; Kunze *et al.* (1985) *J. Basic Microbiol.* 25:141; Candida]; [Gleeson *et al.* (1986) *J. Gen. Microbiol.* 132:3459; Roggenkamp *et al.* (1986) *Mol. Gen. Genet.* 202:302; Hansenula]; [Das *et al.* (1984) *J. Bacteriol.* 158:1165; De Louvencourt *et al.* (1983) *J. Bacteriol.* 154:1165; Van den Berg *et al.* (1990) *BioTechnology* 8:135; Kluyveromyces]; [Cregg *et al.* (1985) *Mol. Cell. Biol.* 5:3376; Kunze *et al.* (1985) *J. Basic Microbiol.* 25:141; US Patent Nos. 4,837,148 and 4,929,555; Pichia]; [Hinnen *et al.* (1978) *Proc. Natl. Acad. Sci. USA* 75:1929; Ito *et al.* (1983) *J. Bacteriol.* 153:163 Saccharomyces]; [Beach and Nurse (1981) *Nature* 300:706; Schizosaccharomyces]; [Davidow *et al.* (1985) *Curr. Genet.* 10:39; Gaillardin *et al.* (1985) *Curr. Genet.* 10:49; Yarrowia].

Antibodies

- 30 As used herein, the term "antibody" refers to a polypeptide or group of polypeptides composed of at least one antibody combining site. An "antibody combining site" is the three-dimensional binding space with an internal surface shape and charge distribution complementary to the features of an epitope of an antigen, which allows a binding of the antibody with the antigen. "Antibody" includes, for example, vertebrate antibodies, hybrid antibodies, chimeric antibodies, humanised antibodies, altered antibodies, univalent antibodies, Fab proteins, and single domain antibodies.

- 35 Antibodies against the proteins of the invention are useful for affinity chromatography, immunoassays, and distinguishing/identifying streptococcus proteins.

Antibodies to the proteins of the invention, both polyclonal and monoclonal, may be prepared by conventional methods. In general, the protein is first used to immunize a suitable animal, preferably a mouse, rat, rabbit or goat. Rabbits and goats are preferred for the preparation of polyclonal sera due to the volume of serum obtainable, and the availability of labeled anti-rabbit and anti-goat antibodies. Immunization is generally performed by mixing or emulsifying the protein in saline, preferably in an adjuvant such as Freund's complete adjuvant, and injecting the mixture or emulsion parenterally (generally subcutaneously or intramuscularly). A dose of 50-200 µg/injection is typically sufficient. Immunization is generally boosted 2-6 weeks later with one or more injections of the protein in saline, preferably using Freund's incomplete adjuvant. One may alternatively generate antibodies *in vitro* immunization using methods known in the art, which for the purposes of this invention is considered equivalent to *in vivo* immunization. Polyclonal antisera is obtained by bleeding the immunized animal into a glass or plastic container, incubating the blood at 25°C for one hour, followed by incubating at 4°C for 2-18 hours. The serum is recovered by centrifugation (eg. 1,000g for 10 minutes). About 20-50 ml per bleed may be obtained from rabbits.

- 40 Monoclonal antibodies are prepared using the standard method of Kohler & Milstein [*Nature* (1975) 256:495-96], or a modification thereof. Typically, a mouse or rat is immunized as described above. However, rather than bleeding the animal to extract serum, the spleen (and optionally several large lymph nodes) is removed and dissociated into single cells. If desired, the
- 50

spleen cells may be screened (after removal of nonspecifically adherent cells) by applying a cell suspension to a plate or well coated with the protein antigen. B-cells expressing membrane-bound immunoglobulin specific for the antigen bind to the plate, and are not rinsed away with the rest of the suspension. Resulting B-cells, or all dissociated spleen cells, are then induced to fuse with myeloma cells to form hybridomas, and are cultured in a selective medium (eg. hypoxanthine, aminopterin, thymidine medium, "HAT"). The resulting hybridomas are plated by limiting dilution, and are assayed for production of antibodies which bind specifically to the immunizing antigen (and which do not bind to unrelated antigens). The selected MAB-secreting hybridomas are then cultured either *in vitro* (eg. in tissue culture bottles or hollow fiber reactors), or *in vivo* (as ascites in mice).

If desired, the antibodies (whether polyclonal or monoclonal) may be labeled using conventional techniques. Suitable labels include fluorophores, chromophores, radioactive atoms (particularly ^{32}P and ^{125}I), electron-dense reagents, enzymes, and ligands having specific binding partners. Enzymes are typically detected by their activity. For example, horseradish peroxidase is usually detected by its ability to convert 3,3',5,5'-tetramethylbenzidine (TMB) to a blue pigment, quantifiable with a spectrophotometer. "Specific binding partner" refers to a protein capable of binding a ligand molecule with high specificity, as for example in the case of an antigen and a monoclonal antibody specific therefor. Other specific binding partners include biotin and avidin or streptavidin, IgG and protein A, and the numerous receptor-ligand couples known in the art. It should be understood that the above description is not meant to categorize the various labels into distinct classes, as the same label may serve in several different modes. For example, ^{125}I may serve as a radioactive label or as an electron-dense reagent. HRP may serve as enzyme or as antigen for a MAb. Further, one may combine various labels for desired effect. For example, MAbs and avidin also require labels in the practice of this invention: thus, one might label a MAb with biotin, and detect its presence with avidin labeled with ^{125}I , or with an anti-biotin MAb labeled with HRP. Other permutations and possibilities will be readily apparent to those of ordinary skill in the art, and are considered as equivalents within the scope of the instant invention.

Pharmaceutical Compositions

Pharmaceutical compositions can comprise either polypeptides, antibodies, or nucleic acid of the invention. The pharmaceutical compositions will comprise a therapeutically effective amount of either polypeptides, antibodies, or polynucleotides of the claimed invention.

The term "therapeutically effective amount" as used herein refers to an amount of a therapeutic agent to treat, ameliorate, or prevent a desired disease or condition, or to exhibit a detectable therapeutic or preventative effect. The effect can be detected by, for example, chemical markers or antigen levels. Therapeutic effects also include reduction in physical symptoms, such as decreased body temperature. The precise effective amount for a subject will depend upon the subject's size and health, the nature and extent of the condition, and the therapeutics or combination of therapeutics selected for administration. Thus, it is not useful to specify an exact effective amount in advance. However, the effective amount for a given situation can be determined by routine experimentation and is within the judgement of the clinician.

For purposes of the present invention, an effective dose will be from about 0.01 mg/kg to 50 mg/kg or 0.05 mg/kg to about 10 mg/kg of the molecule of the invention in the individual to which it is administered.

A pharmaceutical composition can also contain a pharmaceutically acceptable carrier. The term "pharmaceutically acceptable carrier" refers to a carrier for administration of a therapeutic agent, such as antibodies or a polypeptide, genes, and other therapeutic agents. The term refers to any pharmaceutical carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition, and which may be administered without undue toxicity. Suitable carriers may be large, slowly metabolized macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, and inactive virus particles. Such carriers are well known to those of ordinary skill in the art.

Pharmaceutically acceptable salts can be used therein, for example, mineral acid salts such as hydrochlorides, hydrobromides, phosphates, sulfates, and the like; and the salts of organic acids such as acetates, propionates, malonates, benzoates, and the like. A thorough discussion of pharmaceutically acceptable excipients is available in Remington's Pharmaceutical Sciences (Mack Pub. Co., N.J. 1991).

Pharmaceutically acceptable carriers in therapeutic compositions may contain liquids such as water, saline, glycerol and ethanol. Additionally, auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, may be present in such vehicles. Typically, the therapeutic compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection may also be prepared. Liposomes are included within the definition of a pharmaceutically acceptable carrier.

Delivery Methods

Once formulated, the compositions of the invention can be administered directly to the subject. The subjects to be treated can be animals; in particular, human subjects can be treated.

- 5 Direct delivery of the compositions will generally be accomplished by injection, either subcutaneously, intraperitoneally, intravenously or intramuscularly or delivered to the interstitial space of a tissue. The compositions can also be administered into a lesion. Other modes of administration include oral and pulmonary administration, suppositories, and transdermal or transcutaneous applications (eg. see WO98/20734), needles, and gene guns or hypodermic sprays. Dosage treatment may be a single dose schedule or a multiple dose schedule.

Vaccines

- 10 Vaccines according to the invention may either be prophylactic (i.e. to prevent infection) or therapeutic (i.e. to treat disease after infection).

Such vaccines comprise immunising antigen(s), immunogen(s), polypeptide(s), protein(s) or nucleic acid, usually in combination with "pharmaceutically acceptable carriers," which include any carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition. Suitable carriers are typically large, slowly metabolized macromolecules such as proteins, polysaccharides, polyalactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, lipid aggregates (such as oil droplets or liposomes), and inactive virus particles. Such carriers are well known to those of ordinary skill in the art. Additionally, these carriers may function as immunostimulating agents ("adjuvants"). Furthermore, the antigen or immunogen may be conjugated to a bacterial toxoid, such as a toxoid from diphtheria, tetanus, cholera, *H. pylori*, etc. pathogens.

- Preferred adjuvants to enhance effectiveness of the composition include, but are not limited to: (1) oil-in-water emulsion formulations (with or without other specific immunostimulating agents such as muramyl peptides (see below) or bacterial cell wall components), such as for example (a) MF59™ (WO90/14837; Chapter 10 in *Vaccine Design – the subunit and adjuvant approach* (1995) ed. Powell & Newman), containing 5% Squalene, 0.5% Tween 80, and 0.5% Span 85 (optionally containing MTP-PE) formulated into submicron particles using a microfluidizer, (b) SAF, containing 10% Squalene, 0.4% Tween 80, 5% pluronic-blocked polymer L121, and thr-MDP either microfluidized into a submicron emulsion or vortexed to generate a larger particle size emulsion, and (c) Ribi™ adjuvant system (RAS), (Ribi Immunochem, Hamilton, MI) containing 2% Squalene, 0.2% Tween 80, and one or more bacterial cell wall components from the group consisting of monophosphoryl lipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably MPL + CWS (Detox™); (2) saponin adjuvants, such as QS21 or Stimulon™ (Cambridge Bioscience, Worcester, MA) may be used or particles generated therefrom such as ISCOMs (immunostimulating complexes), which ISCOMs may be devoid of additional detergent e.g. WO00/07621; (3) Complete Freund's Adjuvant (CFA) and Incomplete Freund's Adjuvant (IFA); (4) cytokines, such as interleukins (e.g. IL-1, IL-2, IL-4, IL-5, IL-6, IL-7, IL-12 (WO99/44636), etc.), interferons (e.g. gamma interferon), macrophage colony stimulating factor (M-CSF), tumor necrosis factor (TNF), etc.; (5) monophosphoryl lipid A (MPL) or 3-O-deacylated MPL (3dMPL) e.g. GB-2220221, EP-A-0689454; (6) combinations of 3dMPL with, for example, QS21 and/or oil-in-water emulsions e.g. EP-A-0835318, EP-A-0735898, EP-A-0761231; (7) oligonucleotides comprising CpG motifs [Krieg *Vaccine* 2000, 19, 618-622; Krieg *Curr Opin Mol Ther* 2001 3:15-24; Roman *et al.*, *Nat. Med.*, 1997, 3, 849-854; Weiner *et al.*, *PNAS USA*, 1997, 94, 10833-10837; Davis *et al.*, *J. Immunol.*, 1998, 160, 870-876; Chu *et al.*, *J. Exp. Med.*, 1997, 186, 1623-1631; Lipford *et al.*, *Eur. J. Immunol.*, 1997, 27, 2340-2344; Moldoveanu *et al.*, *Vaccine*, 1988, 16, 1216-1224; Krieg *et al.*, *Nature*, 1995, 374, 546-549; Kliman *et al.*, *PNAS USA*, 1996, 93, 2879-2883; Ballas *et al.*, *J. Immunol.*, 1996, 157, 1840-1845; Cowdery *et al.*, *J. Immunol.*, 1996, 156, 4570-4575; Halpern *et al.*, *Cell. Immunol.*, 1996, 167, 72-78; Yamamoto *et al.*, *Jpn. J. Cancer Res.*, 1988, 79, 866-873; Stacey *et al.*, *J. Immunol.*, 1996, 157, 2116-2122; Messina *et al.*, *J. Immunol.*, 1991, 147, 1759-1764; Yi *et al.*, *J. Immunol.*, 1996, 157, 4918-4925; Yi *et al.*, *J. Immunol.*, 1996, 157, 5394-5402; Yi *et al.*, *J. Immunol.*, 1998, 160, 4755-4761; and Yi *et al.*, *J. Immunol.*, 1998, 160, 5898-5906; International patent applications WO96/02555, WO98/16247, WO98/18810, WO98/40100, WO98/55495, WO98/37919 and WO98/52581] i.e. containing at least one CG dinucleotide, with 5-methylcytosine optionally being used in place of cytosine; (8) a polyoxyethylene ether or a polyoxyethylene ester e.g. WO99/52549; (9) a polyoxyethylene sorbitan ester surfactant in combination with an octoxynol (e.g. WO01/21207) or a polyoxyethylene alkyl ether or ester surfactant in combination with at least one additional non-ionic surfactant such as an octoxynol (e.g. WO01/21152); (10) an immunostimulatory oligonucleotide (e.g. a CpG oligonucleotide) and a saponin e.g. WO00/62800; (11) an immunostimulant and a particle of metal salt e.g. WO00/23105; (12) a saponin and an oil-in-water emulsion e.g. WO99/11241; (13) a saponin (e.g. QS21) + 3dMPL + IL-12 (optionally + a sterol) e.g. WO98/57659; (14) aluminium salts, preferably hydroxide or phosphate, but any other suitable salt may also be used (e.g. hydroxyphosphate, oxyhydroxide, orthophosphate, sulphate etc. [e.g. see chapters 8 & 9 of Powell & Newman]). Mixtures of different aluminium

salts may also be used. The salt may take any suitable form (e.g. gel, crystalline, amorphous *etc.*); (15) other substances that act as immunostimulating agents to enhance the efficacy of the composition. Aluminium salts and/or MF59™ are preferred.

As mentioned above, muramyl peptides include, but are not limited to, N-acetyl-muramyl-L-threonine-D-isoglutamine (thr-MDP), N-acetyl-nor-muramyl-L-alanyl-D-isoglutamine (nor-MDP), N-acetyl-muramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-hydroxyphosphoryloxy)-ethylamine (MTP-PE), *etc.*

The immunogenic compositions (e.g. the immunising antigen/immunogen/polypeptide/protein/ nucleic acid, pharmaceutically acceptable carrier, and adjuvant) typically will contain diluents, such as water, saline, glycerol, ethanol, *etc.* Additionally, auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, may be present in such vehicles.

Typically, the immunogenic compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection may also be prepared. The preparation also may be emulsified or encapsulated in liposomes for enhanced adjuvant effect, as discussed above under pharmaceutically acceptable carriers.

Immunogenic compositions used as vaccines comprise an immunologically effective amount of the antigenic or immunogenic polypeptides, as well as any other of the above-mentioned components, as needed. By "immunologically effective amount", it is meant that the administration of that amount to an individual, either in a single dose or as part of a series, is effective for treatment or prevention. This amount varies depending upon the health and physical condition of the individual to be treated, the taxonomic group of individual to be treated (e.g. nonhuman primate, primate, *etc.*), the capacity of the individual's immune system to synthesize antibodies, the degree of protection desired, the formulation of the vaccine, the treating doctor's assessment of the medical situation, and other relevant factors. It is expected that the amount will fall in a relatively broad range that can be determined through routine trials.

The immunogenic compositions are conventionally administered parenterally, e.g. by injection, either subcutaneously, intramuscularly, or transdermally/transcutaneously (e.g. WO98/20734). Additional formulations suitable for other modes of administration include oral and pulmonary formulations, suppositories, and transdermal applications. Dosage treatment may be a single dose schedule or a multiple dose schedule. The vaccine may be administered in conjunction with other immunoregulatory agents.

As an alternative to protein-based vaccines, DNA vaccination may be used [e.g. Robinson & Torres (1997) *Seminars in Immunol* 9:271-283; Donnelly *et al.* (1997) *Annu Rev Immunol* 15:617-648; later herein].

Gene Delivery Vehicles

Gene therapy vehicles for delivery of constructs including a coding sequence of a therapeutic of the invention, to be delivered to the mammal for expression in the mammal, can be administered either locally or systemically. These constructs can utilize viral or non-viral vector approaches in *in vivo* or *ex vivo* modality. Expression of such coding sequence can be induced using endogenous mammalian or heterologous promoters. Expression of the coding sequence *in vivo* can be either constitutive or regulated.

The invention includes gene delivery vehicles capable of expressing the contemplated nucleic acid sequences. The gene delivery vehicle is preferably a viral vector and, more preferably, a retroviral, adenoviral, adeno-associated viral (AAV), herpes viral, or alphavirus vector. The viral vector can also be an astrovirus, coronavirus, orthomyxovirus, papovavirus, paramyxovirus, parvovirus, picornavirus, poxvirus, or togavirus viral vector. See generally, Jolly (1994) *Cancer Gene Therapy* 1:51-64; Kimura (1994) *Human Gene Therapy* 5:845-852; Connelly (1995) *Human Gene Therapy* 6:185-193; and Kaplitt (1994) *Nature Genetics* 6:148-153.

Retroviral vectors are well known in the art and we contemplate that any retroviral gene therapy vector is employable in the invention, including B, C and D type retroviruses, xenotropic retroviruses (for example, NZB-X1, NZB-X2 and NZB9-1 (see O'Neill (1985) *J. Virol.* 53:160) polytropic retroviruses e.g. MCF and MCF-MLV (see Kelly (1983) *J. Virol.* 45:291), spumaviruses and lentiviruses. See RNA Tumor Viruses, Second Edition, Cold Spring Harbor Laboratory, 1985.

Portions of the retroviral gene therapy vector may be derived from different retroviruses. For example, retrovector LTRs may be derived from a Murine Sarcoma Virus, a tRNA binding site from a Rous Sarcoma Virus, a packaging signal from a Murine Leukemia Virus, and an origin of second strand synthesis from an Avian Leukosis Virus.

These recombinant retroviral vectors may be used to generate transduction competent retroviral vector particles by introducing them into appropriate packaging cell lines (see US patent 5,591,624). Retrovirus vectors can be constructed for site-specific integration into host cell DNA by incorporation of a chimeric integrase enzyme into the retroviral particle (see WO96/37626). It is preferable that the recombinant viral vector is a replication defective recombinant virus.

Packaging cell lines suitable for use with the above-described retrovirus vectors are well known in the art, are readily prepared (see WO95/30763 and WO92/05266), and can be used to create producer cell lines (also termed vector cell lines or "VCLs") for the production of recombinant vector particles. Preferably, the packaging cell lines are made from human parent cells (e.g. HT1080 cells) or mink parent cell lines, which eliminates inactivation in human serum.

- 5 Preferred retroviruses for the construction of retroviral gene therapy vectors include Avian Leukosis Virus, Bovine Leukemia Virus, Murine Leukemia Virus, Mink-Cell Focus-Inducing Virus, Murine Sarcoma Virus, Reticuloendotheliosis Virus and Rous Sarcoma Virus. Particularly preferred Murine Leukemia Viruses include 4070A and 1504A (Hartley and Rowe (1976) *J Virol* 19:19-25), Abelson (ATCC No. VR-999), Friend (ATCC No. VR-245), Grafti, Gross (ATCC No. VR-590), Kirsten, Harvey Sarcoma Virus and Rauscher (ATCC No. VR-998) and Moloney Murine Leukemia Virus (ATCC No. VR-190). Such retroviruses may be obtained from depositories or collections such as the American Type Culture Collection ("ATCC") in Rockville, Maryland or isolated from known sources using commonly available techniques.

- Exemplary known retroviral gene therapy vectors employable in this invention include those described in patent applications GB2200651, EP0415731, EP0345242, EP0334301, WO89/02468, WO89/05349, WO89/09271, WO90/02806, WO90/07936, WO94/03622, WO93/25698, WO93/25234, WO93/11230, WO93/10218, WO91/02805, WO91/02825, WO95/07994, US 5,219,740, US 4,405,712, US 4,861,719, US 4,980,289, US 4,777,127, US 5,591,624. See also Vile (1993) *Cancer Res* 53:3860-3864; Vile (1993) *Cancer Res* 53:962-967; Ram (1993) *Cancer Res* 53 (1993) 83-88; Takamiya (1992) *J Neurosci Res* 33:493-503; Baba (1993) *J Neurosurg* 79:729-735; Mann (1983) *Cell* 33:153; Cane (1984) *Proc Natl Acad Sci* 81:6349; and Miller (1990) *Human Gene Therapy* 1.

- Human adenoviral gene therapy vectors are also known in the art and employable in this invention. See, for example, Berkner (1988) *Biotechniques* 6:616 and Rosenfeld (1991) *Science* 252:431, and WO93/07283, WO93/06223, and WO93/07282. Exemplary known adenoviral gene therapy vectors employable in this invention include those described in the above referenced documents and in WO94/12649, WO93/03769, WO93/19191, WO94/28938, WO95/11984, WO95/00655, WO95/27071, WO95/29993, WO95/34671, WO96/05320, WO94/08026, WO94/11506, WO93/06223, WO94/24299, WO95/14102, WO95/24297, WO95/02697, WO94/28152, WO94/24299, WO95/09241, WO95/25807, WO95/05835, WO94/18922 and WO95/09654. Alternatively, administration of DNA linked to killed adenovirus as described in Curiel (1992) *Hum. Gene Ther.* 3:147-154 may be employed. The gene delivery vehicles of the invention also include adenovirus associated virus (AAV) vectors. Leading and preferred examples of such vectors for use in this invention are the AAV-2 based vectors disclosed in Srivastava, WO93/09239. Most preferred AAV vectors comprise the two AAV inverted terminal repeats in which the native Dequences are modified by substitution of nucleotides, such that at least 5 native nucleotides and up to 18 native nucleotides, preferably at least 10 native nucleotides up to 18 native nucleotides, most preferably 10 native nucleotides are retained and the remaining nucleotides of the D-sequence are deleted or replaced with non-native nucleotides. The native D-sequences of the AAV inverted terminal repeats are sequences of 20 consecutive nucleotides in each AAV inverted terminal repeat (e.g. there is one sequence at each end) which are not involved in IIP formation. The non-native replacement nucleotide may be any nucleotide other than the nucleotide found in the native Dequence in the same position. Other employable exemplary AAV vectors are pWP-19, pWN-1, both of which are disclosed in Nahreini (1993) *Gene* 124:257-262. Another example of such an AAV vector is psuh201 (see Samulski (1987) *J. Virol.* 61:3096). Another exemplary AAV vector is the Double-D ITR vector. Construction of the Double-D ITR vector is disclosed in US Patent 5,478,745. Still other vectors are those disclosed in Carter US Patent 4,797,368 and Muzyczka US Patent 5,139,941, Chartejee US Patent 5,474,935, and Kotin WO94/288157. Yet a further example of an AAV vector employable in this invention is SSV9AFABTKneo, which contains the AFP enhancer and albumin promoter and directs expression predominantly in the liver. Its structure and construction are disclosed in Su (1996) *Human Gene Therapy* 7:463-470. Additional AAV gene therapy vectors are described in US 5,354,678, US 5,173,414, US 5,139,941, and US 5,252,479.

- The gene therapy vectors of the invention also include herpes vectors. Leading and preferred examples are herpes simplex virus vectors containing a sequence encoding a thymidine kinase polypeptide such as those disclosed in US 5,288,641 and EP0176170 (Roizman). Additional exemplary herpes simplex virus vectors include HFEM/ICP6-LacZ disclosed in WO95/04139 (Wistar Institute), pHSVlac described in Geller (1988) *Science* 241:1667-1669 and in WO90/09441 and WO92/07945, HSV Us3:pgC-lacZ, described in Fink (1992) *Human Gene Therapy* 3:11-19 and HSV 7134, 2 RH 105 and GAL4 described in EP 0453242 (Breakfield), and those deposited with the ATCC with accession numbers VR-977 and VR-260.

- Also contemplated are alpha virus gene therapy vectors that can be employed in this invention. Preferred alpha virus vectors are Sindbis viruses vectors. Togaviruses, Semliki Forest virus (ATCC VR-67; ATCC VR-1247), Middleberg virus (ATCC

VR-370), Ross River virus (ATCC VR-373; ATCC VR-1246), Venezuelan equine encephalitis virus (ATCC VR923; ATCC VR-1250; ATCC VR-1249; ATCC VR-532), and those described in US patents 5,091,309, 5,217,879, and WO92/10578. More particularly, those alpha virus vectors described in US Serial No. 08/405,627, filed March 15, 1995, WO94/21792, WO92/10578, WO95/07994, US 5,091,309 and US 5,217,879 are employable. Such alpha viruses may be obtained from depositories or collections such as the ATCC in Rockville, Maryland or isolated from known sources using commonly available techniques. Preferably, alphavirus vectors with reduced cytotoxicity are used (see USSN 08/679640).

DNA vector systems such as eukaryotic layered expression systems are also useful for expressing the nucleic acids of the invention. See WO95/07994 for a detailed description of eukaryotic layered expression systems. Preferably, the eukaryotic layered expression systems of the invention are derived from alphavirus vectors and most preferably from Sindbis viral vectors.

- 10 Other viral vectors suitable for use in the present invention include those derived from poliovirus, for example ATCC VR-58 and those described in Evans, *Nature* 339 (1989) 385 and Sabin (1973) *J. Biol. Standardization* 1:115; rhinovirus, for example ATCC VR-1110 and those described in Arnold (1990) *J Cell Biochem* L401; pox viruses such as canary pox virus or vaccinia virus, for example ATCC VR-111 and ATCC VR-2010 and those described in Fisher-Hoch (1989) *Proc Natl Acad Sci* 86:317; Flexner (1989) *Ann NY Acad Sci* 569:86; Flexner (1990) *Vaccine* 8:17; in US 4,603,112 and US 4,769,330 and WO89/01973; SV40 virus, for example ATCC VR-305 and those described in Mulligan (1979) *Nature* 277:108 and Madzak (1992) *J Gen Virol* 73:1533; influenza virus, for example ATCC VR-797 and recombinant influenza viruses made employing reverse genetics techniques as described in US 5,166,057 and in Enami (1990) *Proc Natl Acad Sci* 87:3802-3805; Enami & Palese (1991) *J Virol* 65:2711-2713 and Luytjes (1989) *Cell* 59:110, (see also McMichael (1983) *NEJ Med* 309:13, and Yap (1978) *Nature* 273:238 and *Nature* (1979) 277:108); human immunodeficiency virus as described in EP-0386882 and in Buchschacher (1992) *J. Virol.* 66:2731; measles virus, for example ATCC VR-67 and VR-1247 and those described in EP-0440219; Aura virus, for example ATCC VR-368; Bebaru virus, for example ATCC VR-600 and ATCC VR-1240; Cabassou virus, for example ATCC VR-922; Chikungunya virus, for example ATCC VR-64 and ATCC VR-1241; Fort Morgan Virus, for example ATCC VR-924; Getah virus, for example ATCC VR-369 and ATCC VR-1243; Kyzylagach virus, for example ATCC VR-927; Mayaro virus, for example ATCC VR-66; Mucambo virus, for example ATCC VR-580 and ATCC VR-1244;
- 25 Ndumu virus, for example ATCC VR-371; Pixuna virus, for example ATCC VR-372 and ATCC VR-1245; Tonate virus, for example ATCC VR-925; Tinitit virus, for example ATCC VR-469; Una virus, for example ATCC VR-374; Whataroa virus, for example ATCC VR-926; Y-62-33 virus, for example ATCC VR-375; ONyong virus, Eastern encephalitis virus, for example ATCC VR-65 and ATCC VR-1242; Western encephalitis virus, for example ATCC VR-70, ATCC VR-1251, ATCC VR-622 and ATCC VR-1252; and coronavirus, for example ATCC VR-740 and those described in Hamre (1966) *Proc Soc Exp Biol Med* 121:190.

Delivery of the compositions of this invention into cells is not limited to the above mentioned viral vectors. Other delivery methods and media may be employed such as, for example, nucleic acid expression vectors, polycationic condensed DNA linked or unlinked to killed adenovirus alone, for example see US Serial No. 08/366,787, filed December 30, 1994 and Curiel (1992) *Hum Gene Ther* 3:147-154 ligand linked DNA, for example see Wu (1989) *J Biol Chem* 264:16985-16987, eucaryotic cell delivery vehicles cells, for example see US Serial No.08/240,030, filed May 9, 1994, and US Serial No. 08/404,796, deposition of photopolymerized hydrogel materials, hand-held gene transfer particle gun, as described in US Patent 5,149,655, ionizing radiation as described in US5,206,152 and in WO92/11033, nucleic charge neutralization or fusion with cell membranes. Additional approaches are described in Philip (1994) *Mol Cell Biol* 14:2411-2418 and in Woffendin (1994) *Proc Natl Acad Sci* 91:1581-1585.

- 40 Particle mediated gene transfer may be employed, for example see US Serial No. 60/023,867. Briefly, the sequence can be inserted into conventional vectors that contain conventional control sequences for high level expression, and then incubated with synthetic gene transfer molecules such as polymeric DNA-binding cations like polylysine, protamine, and albumin, linked to cell targeting ligands such as asialoorosomucoid, as described in Wu & Wu (1987) *J. Biol. Chem.* 262:4429-4432, insulin as described in Huckled (1990) *Biochem Pharmacol* 40:253-263, galactose as described in Plank (1992) *Bioconjugate Chem* 3:533-539, lactose or transferrin.

Naked DNA may also be employed. Exemplary naked DNA introduction methods are described in WO 90/11092 and US 5,580,859. Uptake efficiency may be improved using biodegradable latex beads. DNA coated latex beads are efficiently transported into cells after endocytosis initiation by the beads. The method may be improved further by treatment of the beads to increase hydrophobicity and thereby facilitate disruption of the endosome and release of the DNA into the cytoplasm.

- 50 Liposomes that can act as gene delivery vehicles are described in US 5,422,120, WO95/13796, WO94/23697, WO91/14445 and EP-524,968. As described in USSN. 60/023,867, on non-viral delivery, the nucleic acid sequences encoding a polypeptide

can be inserted into conventional vectors that contain conventional control sequences for high level expression, and then be incubated with synthetic gene transfer molecules such as polymeric DNA-binding cations like polylysine, protamine, and albumin, linked to cell targeting ligands such as asialoorosomucoid, insulin, galactose, lactose, or transferrin. Other delivery systems include the use of liposomes to encapsulate DNA comprising the gene under the control of a variety of tissue-specific or ubiquitously-active promoters. Further non-viral delivery suitable for use includes mechanical delivery systems such as the approach described in Woffendin *et al* (1994) *Proc. Natl. Acad. Sci. USA* 91(24):11581-11585. Moreover, the coding sequence and the product of expression of such can be delivered through deposition of photopolymerized hydrogel materials. Other conventional methods for gene delivery that can be used for delivery of the coding sequence include, for example, use of hand-held gene transfer particle gun, as described in US 5,149,655; use of ionizing radiation for activating transferred gene, as described in US 5,206,152 and WO92/11033

Exemplary liposome and polycationic gene delivery vehicles are those described in US 5,422,120 and 4,762,915; in WO 95/13796; WO94/23697; and WO91/14445; in EP-0524968; and in Stryer, *Biochemistry*, pages 236-240 (1975) W.H. Freeman, San Francisco; Szoka (1980) *Biochem Biophys Acta* 600:1; Bayer (1979) *Biochem Biophys Acta* 550:464; Rivnay (1987) *Meth Enzymol* 149:119; Wang (1987) *Proc Natl Acad Sci* 84:7851; Plant (1989) *Anal Biochem* 176:420.

A polynucleotide composition can comprises therapeutically effective amount of a gene therapy vehicle, as the term is defined above. For purposes of the present invention, an effective dose will be from about 0.01 mg/kg to 50 mg/kg or 0.05 mg/kg to about 10 mg/kg of the DNA constructs in the individual to which it is administered.

Delivery Methods

Once formulated, the polynucleotide compositions of the invention can be administered (1) directly to the subject; (2) delivered *ex vivo*, to cells derived from the subject; or (3) *in vitro* for expression of recombinant proteins. The subjects to be treated can be mammals or birds. Also, human subjects can be treated.

Direct delivery of the compositions will generally be accomplished by injection, either subcutaneously, intraperitoneally, intravenously or intramuscularly or delivered to the interstitial space of a tissue. The compositions can also be administered into a lesion. Other modes of administration include oral and pulmonary administration, suppositories, and transdermal or transcutaneous applications (eg. see WO98/20734), needles, and gene guns or hypodermis. Dosage treatment may be a single dose schedule or a multiple dose schedule.

Methods for the *ex vivo* delivery and reimplantation of transformed cells into a subject are known in the art and described in eg. WO93/14778. Examples of cells useful in *ex vivo* applications include, for example, stem cells, particularly hematopoietic, lymph cells, macrophages, dendritic cells, or tumor cells.

Generally, delivery of nucleic acids for both *ex vivo* and *in vitro* applications can be accomplished by the following procedures, for example, dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei, all well known in the art.

Polynucleotide and polypeptide pharmaceutical compositions

In addition to the pharmaceutically acceptable carriers and salts described above, the following additional agents can be used with polynucleotide and/or polypeptide compositions.

A. Polypeptides

One example are polypeptides which include, without limitation: asialoorosomucoid (ASOR); transferrin; asialoglycoproteins; antibodies; antibody fragments; ferritin; interleukins; interferons; granulocyte, macrophage colony stimulating factor (GM-CSF), granulocyte colony stimulating factor (G-CSF), macrophage colony stimulating factor (M-CSF), stem cell factor and erythropoietin. Viral antigens, such as envelope proteins, can also be used. Also, proteins from other invasive organisms, such as the 17 amino acid peptide from the circumsporozoite protein of *Plasmodium falciparum* known as RII.

B. Hormones, Vitamins, etc.

Other groups that can be included are, for example: hormones, steroids, androgens, estrogens, thyroid hormone, or vitamins, folic acid.

C.Polyalkylenes, Polysaccharides, etc.

Also, polyalkylene glycol can be included with the desired polynucleotides/polypeptides. In a preferred embodiment, the polyalkylene glycol is polyethylene glycol. In addition, mono-, di-, or polysaccharides can be included. In a preferred embodiment of this aspect, the polysaccharide is dextran or DEAE-dextran. Also, chitosan and poly(lactide-co-glycolide)

5 D.Lipids, and Liposomes

The desired polynucleotide/polypeptide can also be encapsulated in lipids or packaged in liposomes prior to delivery to the subject or to cells derived therefrom.

Lipid encapsulation is generally accomplished using liposomes which are able to stably bind or entrap and retain nucleic acid. The ratio of condensed polynucleotide to lipid preparation can vary but will generally be around 1:1 (mg DNA:micromoles lipid), or more of lipid. For a review of the use of liposomes as carriers for delivery of nucleic acids, see, Hug and Sleight (1991) *Biochim. Biophys. Acta*. 1097:1-17; Straubinger (1983) *Meth. Enzymol.* 101:512-527.

- 10 Liposomal preparations for use in the present invention include cationic (positively charged), anionic (negatively charged) and neutral preparations. Cationic liposomes have been shown to mediate intracellular delivery of plasmid DNA (Felgner (1987) *Proc. Natl. Acad. Sci. USA* 84:7413-7416; mRNA (Malone (1989) *Proc. Natl. Acad. Sci. USA* 86:6077-6081); and purified transcription factors (Debs (1990) *J. Biol. Chem.* 265:10189-10192), in functional form.

- 15 Cationic liposomes are readily available. For example, N[1-2,3-di(octyloxy)propyl]-N,N,N-trimethylammonium (DOTMA) liposomes are available under the trademark Lipofectin, from GIBCO BRL, Grand Island, NY. (See, also, Felgner *supra*). Other commercially available liposomes include transfectane (DDAB/DOPE) and DOTAP/DOPE (Boehringer). Other cationic liposomes can be prepared from readily available materials using techniques well known in the art. See, eg. Szoka (1978) *Proc. Natl. Acad. Sci. USA* 75:4194-4198; WO90/11092 for a description of the synthesis of DOTAP (1,2-bis(octyloxy)-3-(trimethylammonio)propane) liposomes.

- 20 Similarly, anionic and neutral liposomes are readily available, such as from Avanti Polar Lipids (Birmingham, AL), or can be easily prepared using readily available materials. Such materials include phosphatidyl choline, cholesterol, phosphatidyl ethanolamine, dioleoylphosphatidyl choline (DOPC), dioleoylphosphatidyl glycerol (DOPG), dioleoylphosphatidyl ethanolamine (DOPE), among others. These materials can also be mixed with the DOTMA and DOTAP starting materials in appropriate ratios. Methods for making liposomes using these materials are well known in the art.

- The liposomes can comprise multilamellar vesicles (MLVs), small unilamellar vesicles (SUVs), or large unilamellar vesicles (LUVs). The various liposome-nucleic acid complexes are prepared using methods known in the art. See eg. Straubinger (1983) *Meth. Immunol.* 101:512-527; Szoka (1978) *Proc. Natl. Acad. Sci. USA* 75:4194-4198; Papahadjopoulos (1975) *Biochim. Biophys. Acta* 394:483; Wilson (1979) *Cell* 17:77; Deamer & Bangham (1976) *Biochim. Biophys. Acta* 443:629; Ostro (1977) *Biochem. Biophys. Res. Commun.* 76:836; Fraley (1979) *Proc. Natl. Acad. Sci. USA* 76:3348; Enoch & Strittmatter (1979) *Proc. Natl. Acad. Sci. USA* 76:145; Fraley (1980) *J. Biol. Chem.* (1980) 255:10431; Szoka & Papahadjopoulos (1978) *Proc. Natl. Acad. Sci. USA* 75:145; and Schaefer-Ridder (1982) *Science* 215:166.

E.Lipoproteins

- 35 In addition, lipoproteins can be included with the polynucleotide/polypeptide to be delivered. Examples of lipoproteins to be utilized include: chylomicrons, HDL, IDL, LDL, and VLDL. Mutants, fragments, or fusions of these proteins can also be used. Also, modifications of naturally occurring lipoproteins can be used, such as acetylated LDL. These lipoproteins can target the delivery of polynucleotides to cells expressing lipoprotein receptors. Preferably, if lipoproteins are including with the polynucleotide to be delivered, no other targeting ligand is included in the composition.

- 40 Naturally occurring lipoproteins comprise a lipid and a protein portion. The protein portion are known as apoproteins. At the present, apoproteins A, B, C, D, and E have been isolated and identified. At least two of these contain several proteins, designated by Roman numerals, AI, AII, AIV; CI, CII, CIII.

- A lipoprotein can comprise more than one apoprotein. For example, naturally occurring chylomicrons comprises of A, B, C & E, over time these lipoproteins lose A and acquire C & E. VLDL comprises A, B, C & E apoproteins, LDL comprises apoprotein B; and HDL comprises apoproteins A, C, & E.

The amino acid of these apoproteins are known and are described in, for example, Breslow (1985) *Annu Rev. Biochem.* 54:699; Law (1986) *Adv. Exp. Med. Biol.* 151:162; Chen (1986) *J Biol Chem* 261:12918; Kane (1980) *Proc Natl Acad Sci USA* 77:2465; and Utermann (1984) *Hum Genet* 65:232.

Lipoproteins contain a variety of lipids including, triglycerides, cholesterol (free and esters), and phospholipids. The composition of the lipids varies in naturally occurring lipoproteins. For example, chylomicrons comprise mainly triglycerides. A more detailed description of the lipid content of naturally occurring lipoproteins can be found, for example, in *Meth. Enzymol.* 128 (1986). The composition of the lipids are chosen to aid in conformation of the apoprotein for receptor binding activity. The composition of lipids can also be chosen to facilitate hydrophobic interaction and association with the polynucleotide binding molecule.

Naturally occurring lipoproteins can be isolated from serum by ultracentrifugation, for instance. Such methods are described in *Meth. Enzymol.* (supra); Pitas (1980) *J. Biochem.* 255:5454-5460 and Mahey (1979) *J. Clin. Invest.* 64:743-750. Lipoproteins can also be produced by *in vitro* or recombinant methods by expression of the apoprotein genes in a desired host cell. See, for example, Atkinson (1986) *Annu Rev Biophys Chem* 15:403 and Radding (1958) *Biochim Biophys Acta* 30: 443. Lipoproteins can also be purchased from commercial suppliers, such as Biomedical Technologies, Inc., Stoughton, MA, USA. Further description of lipoproteins can be found in WO98/06437..

F. Polycationic Agents

Polycationic agents can be included, with or without lipoprotein, in a composition with the desired polynucleotide/polypeptide to be delivered.

Polycationic agents, typically, exhibit a net positive charge at physiological relevant pH and are capable of neutralizing the electrical charge of nucleic acids to facilitate delivery to a desired location. These agents have both *in vitro*, *ex vivo*, and *in vivo* applications. Polycationic agents can be used to deliver nucleic acids to a living subject either intramuscularly, subcutaneously, *etc.*

The following are examples of useful polypeptides as polycationic agents: polylysine, polyarginine, polyornithine, and protamine. Other examples include histones, protamines, human serum albumin, DNA binding proteins, non-histone chromosomal proteins, coat proteins from DNA viruses, such as (X174, transcriptional factors also contain domains that bind DNA and therefore may be useful as nucleic acid condensing agents. Briefly, transcriptional factors such as C/EBP, eJun, c-fos, AP-1, AP-2, AP-3, CPF, Prot-1, Sp-1, Oct-1, Oct-2, CREP, and TFIID contain basic domains that bind DNA sequences.

Organic polycationic agents include: spermine, spermidine, and putrescine.

The dimensions and of the physical properties of a polycationic agent can be extrapolated from the list above, to construct other polypeptide polycationic agents or to produce synthetic polycationic agents.

Synthetic polycationic agents which are useful include, for example, DEAB-dextran, polybrene. Lipofectin™, and lipofectAMINE™ are monomers that form polycationic complexes when combined with polynucleotides/polypeptides.

Immunodiagnostic Assays

Streptococcus antigens of the invention can be used in immunoassays to detect antibody levels (or, conversely, anti-streptococcus antibodies can be used to detect antigen levels). Immunoassays based on well defined, recombinant antigens can be developed to replace invasive diagnostics methods. Antibodies to streptococcus proteins within biological samples, including for example, blood or serum samples, can be detected. Design of the immunoassays is subject to a great deal of variation, and a variety of these are known in the art. Protocols for the immunoassay may be based, for example, upon competition, or direct reaction, or sandwich type assays. Protocols may also, for example, use solid supports, or may be by immunoprecipitation. Most assays involve the use of labeled antibody or polypeptide; the labels may be, for example, fluorescent, chemiluminescent, radioactive, or dye molecules. Assays which amplify the signals from the probe are also known; examples of which are assays which utilize biotin and avidin, and enzyme-labeled and mediated immunoassays, such as ELISA assays.

Kits suitable for immunodiagnosis and containing the appropriate labeled reagents are constructed by packaging the appropriate materials, including the compositions of the invention, in suitable containers, along with the remaining reagents and materials (for example, suitable buffers, salt solutions, *etc.*) required for the conduct of the assay, as well as suitable set of assay instructions.

Nucleic Acid Hybridization

"Hybridization" refers to the association of two nucleic acid sequences to one another by hydrogen bonding. Typically, one sequence will be fixed to a solid support and the other will be free in solution. Then, the two sequences will be placed in contact with one another under conditions that favor hydrogen bonding. Factors that affect this bonding include: the type and volume of solvent; reaction temperature; time of hybridization; agitation; factors to block the non-specific attachment of the liquid phase sequence to the solid support (Denhardt's reagent or BLOTTO); concentration of the sequences; use of compounds to increase the rate of association of sequences (dextran sulfate or polyethylene glycol); and the stringency of the washing conditions following hybridization. See Sambrook *et al.* [supra] Volume 2, chapter 9, pages 9.47 to 9.57.

"Stringency" refers to conditions in a hybridization reaction that favor association of very similar sequences over sequences that differ. For example, the combination of temperature and salt concentration should be chosen that is approximately 120 to 200°C below the calculated T_m of the hybrid under study. The temperature and salt conditions can often be determined empirically in preliminary experiments in which samples of genomic DNA immobilized on filters are hybridized to the sequence of interest and then washed under conditions of different stringencies. See Sambrook *et al.* at page 9.50.

Variables to consider when performing, for example, a Southern blot are (1) the complexity of the DNA being blotted and (2) the homology between the probe and the sequences being detected. The total amount of the fragment(s) to be studied can vary a magnitude of 10, from 0.1 to 1 µg for a plasmid or phage digest to 10^3 to 10^6 g for a single copy gene in a highly complex eukaryotic genome. For lower complexity polynucleotides, substantially shorter blotting, hybridization, and exposure times, a smaller amount of starting polynucleotides, and lower specific activity of probes can be used. For example, a single-copy yeast gene can be detected with an exposure time of only 1 hour starting with 1 µg of yeast DNA, blotting for two hours, and hybridizing for 48 hours with a probe of 10^8 cpm/µg. For a single-copy mammalian gene a conservative approach would start with 10 µg of DNA, blot overnight, and hybridize overnight in the presence of 10% dextran sulfate using a probe of greater than 10^8 cpm/µg, resulting in an exposure time of ~24 hours.

Several factors can affect the melting temperature (T_m) of a DNA-DNA hybrid between the probe and the fragment of interest, and consequently, the appropriate conditions for hybridization and washing. In many cases the probe is not 100% homologous to the fragment. Other commonly encountered variables include the length and total G+C content of the hybridizing sequences and the ionic strength and formamide content of the hybridization buffer. The effects of all of these factors can be approximated by a single equation:

$$T_m = 81 + 16.6[\log_{10} C] + 0.41\%(G + C) - 0.6(\%\text{formamide}) - 600/n - 1.5(\%\text{mismatch}).$$

where C is the salt concentration (monovalent ions) and n is the length of the hybrid in base pairs (slightly modified from Meinkoth & Wahl (1984) *Anal. Biochem.* 138: 267-284).

In designing a hybridization experiment, some factors affecting nucleic acid hybridization can be conveniently altered. The temperature of the hybridization and washes and the salt concentration during the washes are the simplest to adjust. As the temperature of the hybridization increases (*i.e.* stringency), it becomes less likely for hybridization to occur between strands that are nonhomologous, and as a result, background decreases. If the radiolabeled probe is not completely homologous with the immobilized fragment (as is frequently the case in gene family and interspecies hybridization experiments), the hybridization temperature must be reduced, and background will increase. The temperature of the washes affects the intensity of the hybridizing band and the degree of background in a similar manner. The stringency of the washes is also increased with decreasing salt concentrations.

In general, convenient hybridization temperatures in the presence of 50% formamide are 42°C for a probe with is 95% to 100% homologous to the target fragment, 37°C for 90% to 95% homology, and 32°C for 85% to 90% homology. For lower homologies, formamide content should be lowered and temperature adjusted accordingly, using the equation above. If the homology between the probe and the target fragment are not known, the simplest approach is to start with both hybridization and wash conditions which are nonstringent. If non-specific bands or high background are observed after autoradiography, the filter can be washed at high stringency and reexposed. If the time required for exposure makes this approach impractical, several hybridization and/or washing stringencies should be tested in parallel.

Nucleic Acid Probe Assays

Methods such as PCR, branched DNA probe assays, or blotting techniques utilizing nucleic acid probes according to the invention can determine the presence of cDNA or mRNA. A probe is said to "hybridize" with a sequence of the invention if it can form a duplex or double stranded complex, which is stable enough to be detected.

The nucleic acid probes will hybridize to the streptococcus nucleotide sequences of the invention (including both sense and antisense strands). Though many different nucleotide sequences will encode the amino acid sequence, the native streptococcus sequence is preferred because it is the actual sequence present in cells. mRNA represents a coding sequence and so a probe should be complementary to the coding sequence; single-stranded cDNA is complementary to mRNA, and so a cDNA probe should be complementary to the non-coding sequence.

The probe sequence need not be identical to the streptococcus sequence (or its complement) — some variation in the sequence and length can lead to increased assay sensitivity if the nucleic acid probe can form a duplex with target nucleotides, which can be detected. Also, the nucleic acid probe can include additional nucleotides to stabilize the formed duplex. Additional streptococcus sequence may also be helpful as a label to detect the formed duplex. For example, a non-complementary nucleotide sequence

may be attached to the 5' end of the probe, with the remainder of the probe sequence being complementary to a streptococcus sequence. Alternatively, non-complementary bases or longer sequences can be interspersed into the probe, provided that the probe sequence has sufficient complementarity with the a streptococcus sequence in order to hybridize therewith and thereby form a duplex which can be detected.

- 5 The exact length and sequence of the probe will depend on the hybridization conditions (e.g. temperature, salt condition etc.). For example, for diagnostic applications, depending on the complexity of the analyte sequence, the nucleic acid probe typically contains at least 10-20 nucleotides, preferably 15-25, and more preferably at least 30 nucleotides, although it may be shorter than this. Short primers generally require cooler temperatures to form sufficiently stable hybrid complexes with the template.

- 10 Probes may be produced by synthetic procedures, such as the triester method of Matteucci *et al.* [*J. Am. Chem. Soc.* (1981) 103:3185], or according to Urdea *et al.* [*Proc. Natl. Acad. Sci. USA* (1983) 80: 7461], or using commercially available automated oligonucleotide synthesizers.

The chemical nature of the probe can be selected according to preference. For certain applications, DNA or RNA are appropriate. For other applications, modifications may be incorporated e.g. backbone modifications, such as phosphorothioates or methylphosphonates, can be used to increase *in vivo* half-life, alter RNA affinity, increase nuclease resistance etc. [eg. see Agrawal & Iyer (1995) *Curr Opin Biotechnol* 6:12-19; Agrawal (1996) *TIBTECH* 14:376-387]; analogues such as peptide nucleic acids may also be used [eg. see Corey (1997) *TIBTECH* 15:224-229; Buchardt *et al.* (1993) *TIBTECH* 11:384-386].

- 15 Alternatively, the polymerase chain reaction (PCR) is another well-known means for detecting small amounts of target nucleic acid. The assay is described in Mullis *et al.* [*Meth. Enzymol.* (1987) 155:335-350] & US patents 4,683,195 & 4,683,202. Two "primer" nucleotides hybridize with the target nucleic acids and are used to prime the reaction. The primers can comprise sequence that does not hybridize to the sequence of the amplification target (or its complement) to aid with duplex stability or, for example, to incorporate a convenient restriction site. Typically, such sequence will flank the desired streptococcus sequence.

- 20 A thermostable polymerase creates copies of target nucleic acids from the primers using the original target nucleic acids as a template. After a threshold amount of target nucleic acids are generated by the polymerase, they can be detected by more traditional methods, such as Southern blots. When using the Southern blot method, the labelled probe will hybridize to the streptococcus sequence (or its complement).

- 25 Also, mRNA or cDNA can be detected by traditional blotting techniques described in Sambrook *et al* [*supra*]. mRNA, or cDNA generated from mRNA using a polymerase enzyme, can be purified and separated using gel electrophoresis. The nucleic acids on the gel are then blotted onto a solid support, such as nitrocellulose. The solid support is exposed to a labelled probe and then washed to remove any unhybridized probe. Next, the duplexes containing the labeled probe are detected. Typically, the probe is labelled with a radioactive moiety.

BRIEF DESCRIPTION OF DRAWINGS

- 30 **Figures 1 to 85, 119 to 188, 238 and 239** show SDS-PAGE analysis of total cell extracts from cultures of recombinant *E.coli* expressing GBS proteins of the invention. Lane 1 in each gel (except for Figure 185) contains molecular weight markers. These are 94, 67, 43, 30, 20.1 & 14.4 kDa (except for Figures 7, 8, 10, 11, 13, 14, 15 and 119-170, which use 250, 150, 100, 75, 50, 37, 25, 15 & 10 kDa).

Figure 86A shows the pDEST15 vector and **Figure 86B** shows the pDEST17-1 vector.

Figures 88 to 118 and 247 to 319 show protein characterisation data for various proteins of the invention.

- 40 **Figures 189 to 237 and 240 to 246** show SDS-PAGE analysis of purified GBS proteins of the invention. The left-hand lane contains molecular weight markers. These are 94, 67, 43, 30, 20.1 & 14.4 kDa.

MODES FOR CARRYING OUT THE INVENTION

The following examples describe nucleic acid sequences which have been identified in *Streptococcus*, along with their inferred translation products. The examples are generally in the following format:

- a nucleotide sequence which has been identified in *Streptococcus*
- 5 • the inferred translation product of this sequence
- a computer analysis (*e.g.* PSORT output) of the translation product, indicating antigenicity

Most examples describe nucleotide sequences from *S.agalactiae*. The specific strain which was sequenced was from serotype V, and is a clinical strain isolated in Italy which expresses the R antigen (ISS/Rome/Italy collection, strain.2603 V/R). For several of these examples, the corresponding
10 sequences from *S.pyogenes* are also given. Where GBS and GAS show homology in this way, there is conservation between species which suggests an essential function and also gives good cross-species reactivity.

In contrast, several examples describe nucleotide sequences from GAS for which no homolog in GBS has been identified. This lack of homology gives molecules which are useful for distinguishing GAS
15 from GBS and for making GAS-specific products. The same is true for GBS sequences which lack GAS homologs *e.g.* these are useful for making GBS-specific products.

The examples typically include details of homology to sequences in the public databases. Proteins that are similar in sequence are generally similar in both structure and function, and the homology often indicates a common evolutionary origin. Comparison with sequences of proteins of known function is
20 widely used as a guide for the assignment of putative protein function to a new sequence and has proved particularly useful in whole-genome analyses.

Various tests can be used to assess the *in vivo* immunogenicity of the proteins identified in the examples. For example, the proteins can be expressed recombinantly and used to screen patient sera by immunoblot. A positive reaction between the protein and patient serum indicates that the patient has
25 previously mounted an immune response to the protein in question *i.e.* the protein is an immunogen. This method can also be used to identify immunodominant proteins. The mouse model used in the examples can also be used.

The recombinant protein can also be conveniently used to prepare antibodies *e.g.* in a mouse. These can be used for direct confirmation that a protein is located on the cell-surface. Labelled antibody (*e.g.*
30 fluorescent labelling for FACS) can be incubated with intact bacteria and the presence of label on the bacterial surface confirms the location of the protein.

For many GBS proteins, the following data are given:

- SDS-PAGE analysis of total recombinant *E.coli* cell extracts for GBS protein expression
- SDS-PAGE analysis after the protein purification

- Western-blot analysis of GBS total cell extract using antisera raised against recombinant proteins
- FACS and ELISA analysis against GBS using antisera raised against recombinant proteins
- Results of the *in vivo* passive protection assay

Details of experimental techniques used are presented below:

5 Sequence analysis

Open reading frames (ORFs) within nucleotide sequences were predicted using the GLIMMER program [Salzberg *et al.* (1998) *Nucleic Acids Res* 26:544-8]. Where necessary, start codons were modified and corrected manually on the basis of the presence of ribosome-binding sites and promoter regions on the upstream DNA sequence.

- 10 ORFs were then screened against the non-redundant protein databases using the programs BLASTp [Altschul *et al.* (1990) *J. Mol. Biol.* 215:403-410] and PRAZE, a modification of the Smith-Waterman algorithm [Smith & Waterman (1981) *J Mol Biol* 147:195-7; see Fleischmann *et al* (1995) *Science* 269:496-512].

- 15 Leader peptides within the ORFs were located using three different approaches: (i) PSORT [Nakai (1991) *Bull. Inst. Chem. Res., Kyoto Univ.* 69:269-291; Horton & Nakai (1996) *Intellig. Syst. Mol. Biol.* 4:109-115; Horton & Nakai (1997) *Intellig. Syst. Mol. Biol.* 5:147-152]; (ii) SignalP [Nielsen & Krogh (1998) in *Proceedings of the Sixth International Conference on Intelligent Systems for Molecular Biology (ISMB 6)*, AAAI Press, Menlo Park, California, pp. 122-130; Nielsen *et al.* (1999) *Protein Engineering* 12:3-9; Nielsen *et al.* (1997). *Int. J. Neural Sys.* 8:581-599]; and (iii) visual inspection of the
- 20 ORF sequences. Where a signal sequences is given a "possible site" value, the value represents the C-terminus residue of the signal peptide *e.g.* a "possible site" of 26 means that the signal sequence consists of amino acids 1-26.

- Lipoprotein-specific signal peptides were located using three different approaches: (i) PSORT [see above]; (ii) the "prokaryotic membrane lipoprotein lipid attachment site" PROSITE motif [Hofmann *et al.* (1999) *Nucleic Acids Res.* 27:215-219; Bucher & Bairoch (1994) in *Proceedings 2nd International Conference on Intelligent Systems for Molecular Biology (ISMB-94)*, AAAI Press, pages 53-61]; and
- 25 (iii) the FINDPATTERNS program available in the GCG Wisconsin Package, using the pattern $(M, L, V) \times \{9, 35\} LxxCxx$.

- Transmembrane domains were located using two approaches: (i) PSORT [see above]; (ii) TopPred [von
- 30 Heijne (1992) *J. Mol. Biol.* 225:487-494].

LPXTG motifs, characteristic of cell-wall attached proteins in Gram-positive bacteria [Fischetti *et al.* (1990) *Mol Microbiol* 4:1603-5] were located with FINDPATTERNS using the pattern $(L, I, V, M, Y, F) P_x (T, A, S, G) (G, N, S, T, A, L)$.

RGD motifs, characteristic of cell-adhesion molecules [D'Souza *et al.* (1991) *Trends Biochem Sci* 16:246-50] were located using FINDPATTERNS.

Enzymes belonging to the glycolytic pathway were also selected as antigens, because these have been found experimentally expressed on the surface of *Streptococci* [e.g. Pancholi & Fischetti (1992) *J Exp Med* 176:415-26; Pancholi & Fischetti (1998) *J Biol Chem* 273:14503-15].

Cloning, expression and purification of proteins

GBS genes were cloned to facilitate expression in *E.coli* as two different types of fusion proteins:

- a) proteins having a hexa-histidine tag at the amino-terminus (His-gbs)
- b) proteins having a GST fusion partner at the amino-terminus (Gst-gbs)

10 Cloning was performed using the Gateway™ technology (Life Technologies), which is based on the site-specific recombination reactions that mediate integration and excision of phage lambda into and from the *E.coli* genome. A single cloning experiment included the following steps:

- 1- Amplification of GBS chromosomal DNA to obtain a PCR product coding for a single ORF flanked by *attB* recombination sites.
- 15 2- Insertion of the PCR product into a pDONR vector (containing *attP* sites) through a BP reaction (*attB* x *attP* sites). This reaction gives a so called 'pEntry' vector, which now contains *attL* sites flanking the insert.
- 3- Insertion of the GBS gene into *E.coli* expression vectors (pDestination vectors, containing *attR* sites) through a LR reaction between pEntry and pDestination plasmids (*attL* x *attR* sites).

20 *A) Chromosomal DNA preparation*

For chromosomal DNA preparation, GBS strain 2603 V/R (Istituto Superiore Sanità, Rome) was grown to exponential phase in 2 litres TH Broth (Difco) at 37°C, harvested by centrifugation, and dissolved in 40 ml TES (50 mM Tris pH 8, 5 mM EDTA pH 8, 20% sucrose). After addition of 2.5 ml lysozyme solution (25 mg/ml in TES) and 0.5 ml mutanolysin (Sigma M-9901, 25000U/ml in H₂O), the suspension
25 was incubated at 37°C for 1 hour. 1 ml RNase (20 mg/ml) and 0.1 ml proteinase K (20 mg/ml) were added and incubation was continued for 30 min. at 37°C.

Cell lysis was obtained by adding 5 ml sarkosyl solution (10% N-laurylsarcosine in 250 mM EDTA pH 8.0), and incubating 1 hour at 37°C with frequent inversion. After sequential extraction with phenol, phenol-chloroform and chloroform, DNA was precipitated with 0.3M sodium acetate pH 5.2 and 2
30 volumes of absolute ethanol. The DNA pellet was rinsed with 70% ethanol and dissolved in TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8). DNA concentration was evaluated by OD₂₆₀.

B) Oligonucleotide design

Synthetic oligonucleotide primers were designed on the basis of the coding sequence of each ORF. The aim was to express the protein's extracellular region. Accordingly, predicted signal peptides were omitted (by deducing the 5' end amplification primer sequence immediately downstream from the predicted leader sequence) and C-terminal cell-wall anchoring regions were removed (e.g. LPXTG motifs and downstream amino acids). Where additional nucleotides have been deleted, this is indicated by the suffix 'd' (e.g. 'GBS352d' – see Table V). Conversely, a suffix 'L' refers to expression without these deletions. Deletions of C- or N-terminal residues were also sometimes made, as indicated by a 'C' or 'N' suffix.

- 10 The amino acid sequences of the expressed GBS proteins (including 'd' and 'L' forms *etc.*) are definitively defined by the sequences of the oligonucleotide primers given in Table II.

5' tails of forward primers and 3' tails of reverse primers included *attB1* and *attB2* sites respectively:

Forward primers: 5'-GGGGACAACTTTGTACAAAAAGCAGGCTCT-ORF in frame-3' (the TCT sequence preceding the ORF was omitted when the ORF's first coding triplet began with T).

- 15 **Reverse primers:** 5'-GGGGACCACTTTGTACAAGAAAGCTGGTT-ORF reverse complement-3'.

The number of nucleotides which hybridized to the sequence to be amplified depended on the melting temperature of the primers, which was determined as described by Breslauer *et al.* [*PNAS USA* (1986) 83:3746-50]. The average melting temperature of the selected oligos was 50-55°C for the hybridizing region and 80-85°C for the whole oligos.

20 C) Amplification

The standard PCR protocol was as follows: 50 ng genomic DNA were used as template in the presence of 0.5 µM each primer, 200 µM each dNTP, 1.5 mM MgCl₂, 1x buffer minus Mg⁺⁺ (Gibco-BRL) and 2 units of Taq DNA polymerase (Platinum Taq, Gibco-BRL) in a final volume of 100 µl. Each sample underwent a double-step of amplification: 5 cycles performed using as the hybridizing temperature 50°C, followed by 25 cycles at 68°C.

The standard cycles were as follows:

- | | |
|-------------|--|
| | Denaturation: 94°C, 2 min |
| 5 cycles: | Denaturation: 94°C, 30 seconds |
| | Hybridization: 50°C, 50 seconds |
| 30 | Elongation: 72°C, 1 min. or 2 min. and 40 sec. |
| 25 cycles : | Denaturation: 94°C, 30 seconds |
| | Hybridization: 68°C, 50 seconds |

Elongation: 72°C, 1 min. or 2 min. and 40 sec.

Elongation time was 1 minute for ORFs shorter than 2000bp and 2:40 minutes for ORFs longer than 2000bp. Amplifications were performed using a Gene Amp PCR system 9600 (Perkin Elmer).

- 5 To check amplification results, 2 μ l of each PCR product were loaded onto 1-1.5 agarose gel and the size of amplified fragments was compared with DNA molecular weight standards (DNA marker IX Roche, 1kb DNA ladder Biolabs).

- Single band PCR products were purified by PEG precipitation: 300 μ l of TE buffer and 200 μ l of 30% PEG 8000/30 mM MgCl₂ were added to 100 μ l PCR reaction. After vortexing, the DNA was centrifuged for 20 min at 10000g, washed with 1 vol. 70% ethanol and the pellet dissolved in 30 μ l TE. PCR products smaller than 350 bp were purified using a PCR purification Kit (Qiagen) and eluted with 30 μ l of the provided elution buffer.

In order to evaluate the yield, 2 μ l of the purified DNA were subjected to agarose gel electrophoresis and compared to titrated molecular weight standards.

D) Cloning of PCR products into expression vectors

- 15 Cloning was performed following the GatewayTM technology's "one-tube protocol", which consists of a two step reaction (BP and LR) for direct insertion of PCR products into expression vectors.

BP reaction (*attB* x *attP* sites): The reaction allowed insertion of the PCR product into a pDONR vector. The pDONRTM 201 vector we used contains the killer toxin gene *ccdB* between *attP1* and *attP2* sites to minimize background colonies lacking the PCR insert, and a selectable marker gene for kanamycin resistance. The reaction resulted in a so called pEntry vector, in which the GBS gene was located between *attL1* and *attL2* sites.

60 fmol of PCR product and 100 ng of pDONRTM 201 vector were incubated with 2.5 μ l of BP clonaseTM in a final volume of 12.5 μ l for 4 hours at 25°C.

- 25 **LR reaction** (*attL* x *attR* sites): The reaction allowed the insertion of the GBS gene, now present in the pEntry vector, into *E.coli* expression vectors (pDestination vectors, containing *attR* sites). Two pDestination vectors were used (pDEST15 for N-terminal GST fusions – Figure 86; and pDEST17-1 for N-terminal His-tagged fusions – Figure 87). Both allow transcription of the ORF fusion coding mRNA under T7 RNA polymerase promoter [Studier *et al* (1990) *Meth. Enzymol* 185: 60ff].

- 30 To 5 μ l of BP reaction were added 0.25 μ l of 0.75 M NaCl, 100 ng of destination vector and 1.5 μ l of LR clonaseTM. The reaction was incubated at 25°C for 2 hours and stopped with 1 μ l of 1 mg/ml proteinase K solution at 37°C for 15 min.

1 μ l of the completed reaction was used to transform 50 μ l electrocompetent BL21-SITM cells (0.1 cm, 200 ohms, 25 μ F). BL21-SI cells contain an integrated T7 RNA polymerase gene under the control of the salt-inducible *prU* promoter [Gowrishankar (1985) *J. Bacteriol.* 164:434ff]. After electroporation cells were diluted in 1ml SOC medium (20 g/l bacto-tryptone, 5 g/l yeast extract, 0.58 g/l NaCl, 0.186 g/l KCl, 20 mM glucose, 10 mM MgCl₂) and incubated at 37°C for 1 hour. 200 μ l cells were plated onto LBON plates (Luria Broth medium without NaCl) containing 100 μ g/ ml ampicillin. Plates were then incubated for 16 hours at 37°C.

Entry clones: In order to allow the future preparation of Gateway compatible pEntry plasmids containing genes which might turn out of interest after immunological assays, 2.5 μ l of BP reaction were incubated for 15 min in the presence of 3 μ l 0.15 mg/ml proteinase K solution and then kept at -20°C. The reaction was in this way available to transform *E.coli* competent cells so as to produce Entry clones for future introduction of the genes in other Destination vectors.

E) Protein expression

Single colonies derived from the transformation of LR reactions were inoculated as small-scale cultures in 3 ml LBON 100 μ g/ml ampicillin for overnight growth at 25°C. 50-200 μ l of the culture was inoculated in 3 ml LBON/Amp to an initial OD₆₀₀ of 0.1. The cultures were grown at 37°C until OD₆₀₀ 0.4-0.6 and recombinant protein expression was induced by adding NaCl to a final concentration of 0.3 M. After 2 hour incubation the final OD was checked and the cultures were cooled on ice. 0.5 OD₆₀₀ of cells were harvested by centrifugation. The cell pellet was suspended in 50 μ l of protein Loading Sample Buffer (50 mM TRIS-HCl pH 6.8, 0.5% w/v SDS, 2.5% v/v glycerol, 0.05% w/v Bromophenol Blue, 100 mM DTT) and incubated at 100 °C for 5 min. 10 μ l of sample was analyzed by SDS-PAGE and Coomassie Blue staining to verify the presence of induced protein band.

F) Purification of the recombinant proteins

Single colonies were inoculated in 25 ml LBON 100 μ g/ml ampicillin and grown at 25°C overnight. The overnight culture was inoculated in 500 ml LBON/amp and grown under shaking at 25 °C until OD₆₀₀ values of 0.4-0.6. Protein expression was then induced by adding NaCl to a final concentration of 0.3 M. After 3 hours incubation at 25 °C the final OD₆₀₀ was checked and the cultures were cooled on ice. After centrifugation at 6000 rpm (JA10 rotor, Beckman) for 20 min., the cell pellet was processed for purification or frozen at -20 °C.

Proteins were purified in 1 of 3 ways depending on the fusion partner and the protein's solubility:

Purification of soluble His-tagged proteins from *E.coli*

1. Transfer pellets from -20°C to ice bath and reconstitute each pellet with 10 ml B-PERTM solution (Bacterial-Protein Extraction Reagent, Pierce cat. 78266), 10 μ l of a 100 mM MgCl₂ solution, 50

- μl of DNase I (Sigma D-4263, 100 Kunits in PBS) and 100 μl of 100 mg/ml lysozyme in PBS (Sigma L-7651, final concentration 1 mg/ml).
2. Transfer resuspended pellets in 50 ml centrifuge tubes and leave at room temperature for 30-40 minutes, vortexing 3-4 times.
 - 5 3. Centrifuge 15-20 minutes at about 30-40000 x g.
 4. Prepare Poly-Prep (Bio-Rad) columns containing 1 ml of Fast Flow Ni-activated Chelating Sepharose (Pharmacia). Equilibrate with 50 mM phosphate buffer, 300 mM NaCl, pH 8.0.
 5. Store the pellet at -20°C, and load the supernatant on to the columns.
 6. Discard the flow through.
 - 10 7. Wash with 10 ml 20 mM imidazole buffer, 50 mM phosphate, 300 mM NaCl, pH 8.0.
 8. Elute the proteins bound to the columns with 4.5 ml (1.5 ml + 1.5 ml + 1.5 ml) 250 mM imidazole buffer, 50 mM phosphate, 300 mM NaCl, pH 8.0 and collect three fractions of ~1.5 ml each. Add to each tube 15 μl DTT 200 mM (final concentration 2 mM).
 9. Measure the protein concentration of the collected fractions with the Bradford method and analyse the proteins by SDS-PAGE.
 - 15 10. Store the collected fractions at +4°C while waiting for the results of the SDS-PAGE analysis.
 11. For immunisation prepare 4-5 aliquots of 20-100 μg each in 0.5 ml in 40% glycerol. The dilution buffer is the above elution buffer, plus 2 mM DTT. Store the aliquots at -20°C until immunisation.

Purification of His-tagged proteins from inclusion bodies

- 20 1. Bacteria are collected from 500 ml cultures by centrifugation. If required store bacterial pellets at -20°C. Transfer the pellets from -20°C to room temperature and reconstitute each pellet with 10 ml B-Per™ solution, 10 μl of a 100 mM MgCl₂ solution (final 1 mM), 50 μl of DNase I equivalent to 100 Kunits units in PBS and 100 μl of a 100 mg/ml lysozyme (Sigma L-7651) solution in PBS (equivalent to 10 mg, final concentration 1 mg/ml).
- 25 2. Transfer the resuspended pellets in 50 ml centrifuge tubes and let at room temperature for 30-40 minutes, vortexing 3-4 times.
3. Centrifuge 15 minutes at 30-4000 x g and collect the pellets.
4. Dissolve the pellets with 50 mM TRIS-HCl, 1 mM TCEP {Tris(2-carboxyethyl)-phosphine hydrochloride, Pierce}, 6M guanidine hydrochloride, pH 8.5. Stir for ~ 10 min. with a magnetic bar.
- 30 5. Centrifuge as described above, and collect the supernatant.
6. Prepare Poly-Prep (Bio-Rad) columns containing 1 ml of Fast Flow Ni-activated Chelating Sepharose (Pharmacia). Wash the columns twice with 5 ml of H₂O and equilibrate with 50 mM TRIS-HCl, 1 mM TCEP, 6M guanidine hydrochloride, pH 8.5.

7. Load the supernatants from step 5 onto the columns, and wash with 5 ml of 50 mM TRIS-HCl buffer, 1 mM TCEP, 6M urea, pH 8.5
8. Wash the columns with 10 ml of 20 mM imidazole, 50 mM TRIS-HCl, 6M urea, 1 mM TCEP, pH 8.5. Collect and set aside the first 5 ml for possible further controls.
- 5 9. Elute proteins bound to columns with 4.5ml buffer containing 250 mM imidazole, 50 mM TRIS-HCl, 6M urea, 1 mM TCEP, pH 8.5. Add the elution buffer in three 1.5 ml aliquots, and collect the corresponding three fractions. Add to each fraction 15 μ l DTT (final concentration 2 mM).
- 10 10. Measure eluted protein concentration with Bradford method and analyse proteins by SDS-PAGE.
11. Dialyse overnight the selected fraction against 50 mM Na phosphate buffer, pH 8.8, containing 10% glycerol, 0.5 M arginine, 5 mM reduced glutathione, 0.5 mM oxidized glutathione, 2 M urea.
12. Dialyse against 50 mM Na phosphate buffer, pH 8.8, containing 10% glycerol, 0.5 M arginine, 5 mM reduced glutathione, 0.5 mM oxidized glutathione.
13. Clarify the dialysed protein preparation by centrifugation and discard the non-soluble material and measure the protein concentration with the Bradford method.
- 15 14. For each protein destined to the immunization prepare 4-5 aliquot of 20-100 μ g each in 0.5 ml after having adjusted the glycerol content up to 40%. Store the prepared aliquots at -20° C until immunization.

Purification of GST-fusion proteins from *E.coli*

- 20 1. Bacteria are collected from 500 ml cultures by centrifugation. If required store bacterial pellets at -20° C. Transfer the pellets from -20° C to room temperature and reconstitute each pellet with 10 ml B-PER™ solution, 10 μ l of a 100 mM $MgCl_2$ solution (final 1 mM), 50 μ l of DNase I equivalent to 100 Kunits units in PBS and 100 μ l of a 100 mg/ml lysozyme (Sigma L-7651) solution in PBS (equivalent to 10 mg, final concentration 1 mg/ml).
- 25 2. Transfer the resuspended pellets in 50 ml centrifuge tubes and let at room temperature for 30-40 minutes, vortexing 3-4 times.
3. Centrifuge 15-20 minutes at about 30-40000 x g.
4. Discard centrifugation pellets and load supernatants onto the chromatography columns, as follows.
5. Prepare Poly-Prep (Bio-Rad) columns containing 0.5 ml of Glutathione-Sepharose 4B resin. Wash the columns twice with 1 ml of H_2O and equilibrate with 10 ml PBS, pH 7.4.
- 30 6. Load supernatants on to the columns and discard the flow through.
7. Wash the columns with 10 ml PBS, pH 7.4.
8. Elute proteins bound to columns with 4.5 ml of 50 mM TRIS buffer, 10 mM reduced glutathione, pH 8.0, adding 1.5 ml + 1.5 ml + 1.5 ml and collecting the respective 3 fractions of ~1.5 ml each.

9. Measure protein concentration of the fractions with the Bradford method and analyse the proteins by SDS-PAGE.
10. Store the collected fractions at +4°C while waiting for the results of the SDS-PAGE analysis.
11. For each protein destined for immunisation prepare 4-5 aliquots of 20-100 µg each in 0.5 ml of 40% glycerol. The dilution buffer is 50 mM TRIS-HCl, 2 mM DTT, pH 8.0. Store the aliquots at -20°C until immunisation.

Figures 167 to 170 and 238 to 239

For the experiments shown in Figures 167 to 170, Figure 238 and lanes 2-6 of Figure 239, the GBS proteins were fused at the N-terminus to thioredoxin and at C-terminus to a poly-His tail. The plasmid used for cloning is pBAD-DEST49 (Invitrogen Gateway™ technology) and expression is under the control of an L(+)-Arabinose dependent promoter. For the production of these GBS antigens, bacteria are grown on RM medium (6g/l Na₂HPO₄, 3g/l KH₂PO₄, 0.5 g/l NaCl, 1 g/l NH₄Cl, pH7.4, 2% casaminoacids, 0.2 % glucose, 1 mM MgCl₂) containing 100 µg/ml ampicillin. After incubation at 37°C until cells reach OD₆₀₀=0.5, protein expression is induced by adding 0.2% (v/v) L(+)-Arabinose for 3 hours.

Immunisations with GBS proteins

The purified proteins were used to immunise groups of four CD-1 mice intraperitoneally. 20 µg of each purified protein was injected in Freund's adjuvant at days 1, 21 & 35. Immune responses were monitored by using samples taken on day 0 & 49. Sera were analysed as pools of sera from each group of mice.

FACScan bacteria Binding Assay procedure.

GBS serotype V 2603 V/R strain was plated on TSA blood agar plates and incubated overnight at 37°C. Bacterial colonies were collected from the plates using a sterile dracon swab and inoculated into 100ml Todd Hewitt Broth. Bacterial growth was monitored every 30 minutes by following OD₆₀₀. Bacteria were grown until OD₆₀₀ = 0.7-0.8. The culture was centrifuged for 20 minutes at 5000rpm. The supernatant was discarded and bacteria were washed once with PBS, resuspended in ½ culture volume of PBS containing 0.05% paraformaldehyde, and incubated for 1 hour at 37°C and then overnight at 4°C.

50µl bacterial cells (OD₆₀₀ 0.1) were washed once with PBS and resuspended in 20µl blocking serum (Newborn Calf Serum, Sigma) and incubated for 20 minutes at room temperature. The cells were then incubated with 100µl diluted sera (1:200) in dilution buffer (20% Newborn Calf Serum 0.1% BSA in PBS) for 1 hour at 4°C. Cells were centrifuged at 5000rpm, the supernatant aspirated and cells washed by adding 200µl washing buffer (0.1% BSA in PBS). 50µl R-Phicoerytrin conjugated F(ab)₂ goat anti-mouse, diluted 1:100 in dilution buffer, was added to each sample and incubated for 1 hour at 4°C. Cells were spun down by centrifugation at 5000rpm and washed by adding 200µl of washing buffer. The

supernatant was aspirated and cells resuspended in 200 μ l PBS. Samples were transferred to FACScan tubes and read. The condition for FACScan setting were: FL2 on; FSC-H threshold:54; FSC PMT Voltage: E 02; SSC PMT: 516; Amp. Gains 2.63; FL-2 PMT: 728. Compensation values: 0.

Samples were considered as positive if they had a Δ mean values > 50 channel values.

5 *Whole Extracts preparation*

GBS serotype III COH1 strain and serotype V 2603 V/R strain cells were grown overnight in Todd Hewitt Broth. 1ml of the culture was inoculated into 100ml Todd Hewitt Broth. Bacterial growth was monitored every 30 minutes by following OD₆₀₀. The bacteria were grown until the OD reached 0.7-0.8. The culture was centrifuged for 20 minutes at 5000 rpm. The supernatant was discarded and bacteria
 10 were washed once with PBS, resuspended in 2ml 50mM Tris-HCl, pH 6.8 adding 400 units of Mutanolysin (Sigma-Aldrich) and incubated 3 hrs at 37°C. After 3 cycles of freeze/thaw, cellular debris were removed by centrifugation at 14000g for 15 minutes and the protein concentration of the supernatant was measured by the Bio-Rad Protein assay, using BSA as a standard.

Western blotting

15 Purified proteins (50ng) and total cell extracts (25 μ g) derived from GBS serotype III COH1 strain and serotype V 2603 V/R strain were loaded on 12% or 15% SDS-PAGE and transferred to a nitrocellulose membrane. The transfer was performed for 1 hours at 100V at 4°C, in transferring buffer (25mM Tris base, 192mM glycine, 20% methanol). The membrane was saturated by overnight incubation at 4°C in saturation buffer (5 % skimmed milk, 0.1% Tween 20 in PBS). The membrane was incubated for 1 hour
 20 at room temperature with 1:1000 mouse sera diluted in saturation buffer. The membrane was washed twice with washing buffer (3 % skimmed milk, 0.1% Tween 20 in PBS) and incubated for 1 hour with a 1:5000 dilution of horseradish peroxidase labelled anti-mouse Ig (Bio-Rad). The membrane was washed twice with 0.1% Tween 20 in PBS and developed with the Opti-4CN Substrate Kit (Bio-Rad). The reaction was stopped by adding water.

25 Unless otherwise indicated, lanes 1, 2 and 3 of blots in the drawings are: (1) the purified protein; (2) GBS-III extracts; and (3) GBS-V extracts. Molecular weight markers are also shown.

In vivo passive protection assay in neonatal sepsis mouse model.

The immune sera collected from the CD1 immunized mice were tested in a mouse neonatal sepsis model to verify their protective efficacy in mice challenged with GBS serotype III. Newborn Balb/C littermates
 30 were randomly divided in two groups within 24 hrs from birth and injected subcutaneously with 25 μ l of diluted sera (1:15) from immunized CD1 adult mice. One group received preimmune sera, the other received immune sera. Four hours later all pups were challenged with a 75% lethal dose of the GBS serotype III COH1 strain. The challenge dose obtained diluting a mid log phase culture was administered subcutaneously in 25 μ l of saline. The number of pups surviving GBS infection was assessed every 12
 35 hours for 4 days. Results are in Table III.

Example 1

A DNA sequence (GBSx1402) was identified in *S.agalactiae* <SEQ ID 1> which encodes the amino acid sequence <SEQ ID 2>. Analysis of this protein sequence reveals the following:

```

5      Possible site: 27
      >>> Seems to have an uncleavable N-term signal seq
      INTEGRAL Likelihood = -0.48  Transmembrane 169 - 185 ( 169 - 185)

      ----- Final Results -----
      bacterial membrane -- Certainty=0.1192(Affirmative) < succ>
      bacterial outside -- Certainty=0.0000(Not Clear) < succ>
10     bacterial cytoplasm -- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database.

```

>GP:CA86823 GB:AL353012 hypothetical serine-rich repeat protein
|Schizosaccharomyces pombe|
Identities = 41/152 (26%), Positives = 75/152 (48%), Gaps = 4/152 (2%)

Query: 22 SSIGVADTSDKNDTDSVTTLTKSRKKSELDQSGSTGSSSENSESSESSEPEINPSTNPPT 81
      +++ ++++++ E S+ D S++ SSSS+ESSS ++ S++ +
Sbjct: 132 SSDSESSSEDGDGDESSSDSSSSGGSSGGSSSSSSSSSSSESDNDSSESSDGS 191

Query: 82 TSPSQPSSEENKPDGRKTKE---IGNNNISGGTKLISEDSKFNKASSDQEEVD 138
      S+ S+ D D+++ ++ SS SED+ +S+S+E E D
Sbjct: 192 ESSSDDESSSDGEESSESGSDSSSSSSSESSSESSSENDSSSSSDSESSSED 251

Query: 139 ESSSSKANDGK-KGESKPKFKLPKTGDSHDT 169
      SSS ++D ++ SK ++ DS D+
Sbjct: 252 SDSSSSSSDGTGSSSSKKGSDSSSSSSSDSD 283

```

A related GBS gene <SEQ ID 8785> and protein <SEQ ID 8786> were also identified. Analysis of this protein sequence reveals the following:

```

Lipop: Possible site: -1      Crend: 5
McG: Discrim Score:         6.72
35 GVH: Signal Score (-7.5): -4.34
    Possible site: 27
    >>> Seems to have an uncleavable N-term signal seq
    ALOM program count: 1 value: -0.48 threshold: 0.0
    INTEGRAL Likelihood = -0.48 Transmembrane 169 - 185 (169 - 185)
40 PERIPHERAL Likelihood = 0.16      7
    modified ALOM score: 0.60

*** Reasoning Step: 3

45 ----- Final Results -----
    bacterial membrane --- Certainty=0.1192 (Affirmative) < succ>
    bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
    bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

50 LPXTG motif: 159-163

```

SEQ ID 2 (GBS4) was expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 9 (lane 3; MW 43.1kDa) and Figure 63 (lane 4; MW 50kDa). It was also expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 12 (lane 7; MW 30kDa), Figure 63 (lane 3; MW 30kDa) and in Figure 178 (lane 3; MW 30kDa).

GBS4-GST was purified as shown in Figure 190 (lane 6) and Figure 209 (lane 8).

Purified GBS4-His is shown in Figures 89A, 191 (lane 10), 209 (lane 7) and 228 (lanes 9 & 10).

The purified GBS4-His fusion product was used to immunise mice (lane 2 product; 20g/mouse). The resulting antiserum was used for Western blot (Figure 89B), FACS, and in the *in vivo* passive protection assay (Table III). These tests confirm that the protein is immunoaccessible on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 2

A DNA sequence (GBSx1100) was identified in *S. agalactiae* <SEQ ID 3> which encodes the amino acid sequence <SEQ ID 4>. This protein is predicted to be aggregation promoting protein. Analysis of this protein sequence reveals the following:

```

Possible site: 33
>>> Seems to have a cleavable N-term signal seq.

----- Final Results -----
      bacterial outside --- Certainty=0.3000(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database.

>GP:CAA69725 GB:Y08498 aggregation promoting protein [Lactobacillus gasserii]
Identities = 56/103 (54%), Positives = 69/103 (66%), Gaps = 5/103 (4%)

Query: 82 TASQAEAKGQPT-----IENSMNSSNLSSSDSAAKEETARRESNGSYTAQNQYGYRYQ 136
      T S A A Q T + + + N S S + AAK +A RES G Y+A NQGY G+YQ
Sbjct: 195 TYSYASAGKQTQTTVAQKTTQTTTSTYTNASGSKAAKAMAGRESGPGPYAGNQGYQYIGYQ 254

Query: 137 LSQSYLNGDLSPENQKKVADNYVVSRYGWSAALSFWNSNGWY 179
      LS SYL GD S NQE+VADNYV SKYGS+ A FW +NGWY
Sbjct: 255 LSASYLGGDYSAANQERVADNYVSKRYGWSWTAQKFWQINGWY 297
  
```

No corresponding DNA sequence was identified in *S. pyogenes*.

A related GBS gene <SEQ ID 8709> and protein <SEQ ID 8710> were also identified. Analysis of this protein sequence reveals the following:

```

Lipop: Possible site: -1 Crend: 9
MoG: Discrim Score: 2.59
GVH: Signal Score (-7.5): -0.42
Possible site: 33
>>> Seems to have a cleavable N-term signal seq.
ALOM program count: 0 value: 6.79 threshold: 0.0
PERIPHERAL Likelihood = 6.79 59
modified ALOM score: -1.86

*** Reasoning Step: 3

----- Final Results -----
      bacterial outside --- Certainty=0.3000(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>
  
```

The protein has homology with the following sequences in the databases:

```

57.5/71.3% over 92aa
Lactobacillus gasserii
  
```

EGAD[154417] aggregation promoting protein Insert characterized
 GP[1619598]emb|CAA69725.1||Y08498 aggregation promoting protein Insert characterized

5 ORF01056(547 - 837 of 1137)
 EGAD[154417]164788(205 - 297 of 297) aggregation promoting protein {Lactobacillus
 gasserii}GP[1619598]emb|CAA69725.1||Y08498 aggregat
 ion promoting protein {Lactobacillus gasserii}
 %Match = 14.6
 %Identity = 57.4 %Similarity = 71.3
 10 Matches = 54 Mismatches = 26 Conservative Sub.s = 13

507	537	567	597	627	657	687	717
SENISINADVISIGDVLKLDNSTASQAFKASQPTIENSMNSSLSSSDSAAKEIARRESNGSYTAQNEQYYGRYQLSQ							
:: :	:	:	::	::	:	:	:
15 NVQRTYSAPFVQQRITYSASAKQITTVQAQKTVITTSYTLNAGS---	SEAAAKAWNAGRESGGPYSAQNGSQYIKYQLSA						
	200	210	220	230	240	250	

747	777	807	837	867	897	927	957
SYLNGDLSPECNCKVADNYVVSRYGWSKSAALSPWNSNGWY**KLIKQKRDLLKIKSLCNIPNYSIAR*QIKYNIGNNMR							
	:	:		:			
20 SYLGGDYSAANGERVADNYVKSRYGWSWTGAKQFWQINWY							
	270	280	290				

A related GBS gene <SEQ ID 8711> and protein <SEQ ID 8712> were also identified. Analysis of this
 25 protein sequence reveals the following:

Lipop: Possible site: -1 Crend: 9
 MoG: Discrim Score: 2.59
 GVH: Signal Score (-7.5): -0.42
 Possible site: 33
 30 >>> Seems to have a cleavable N-term signal seq.
 ALOM program count: 0 value: 6.79 threshold: 0.0
 PERIPHERAL Likelihood = 6.79 59
 modified ALOM score: -1.86

35 *** Reasoning Step: 3

----- Final Results -----

bacterial outside --- Certainty=0.3000(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
40 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the databases:

44.0/62.0% over 115aa

Bacillus subtilis

45 EGAD[108478] hypothetical protein Insert characterized OMNI|NT01BS1100 p60-related
 protein Insert characterized
 GP[2226145]emb|CAA74437.1||Y14079 hypothetical protein Insert characterized
 GP[2633272]emb|CAB12776.1||Z99109 similar to cell wall-binding protein Insert
 characterized
 50 PIR[B69825]B69825 cell wall-binding protein homolog yhdD - Insert characterized

ORF01746(340 - 633 of 954)
 EGAD[108478]BS0936(57 - 172 of 488) hypothetical protein {Bacillus subtilis}OMNI|NT01BS1100
 p60-related proteinGP[2226145]emb|CAA74437.1||Y14079 hypothetical protein {Bacillus
 55 subtilis}GP[2633272]emb|CAB12776.1||Z99109 similar to cell wall-binding protein {Bacillus
 subtilis}PIR[B69825]B69825 cell wall-binding protein homolog yhdD - Bacillus subtilis
 %Match = 9.0
 %Identity = 44.0 %Similarity = 62.0
 60 Matches = 44 Mismatches = 35 Conservative Sub.s = 18

120	150	180	210	240	270	300	330
*DQPMVLAFSP**CEKIANFT*RLKIVFWRPFL*FTYIL**ISSKAKQLVIFTRYDSTRIN**KRAYIMISITSVKSK							

MKKKLAAIGLTSATVGTTLVVTFAEAATIKVKSGDSGLWKLQATYNTSVAALTS

65 10 20 30 40 50

```

360      390      435      465      495      525
PFKGLVAGLIVGASIALPLSVSAAS-----YTVKSGDTLSAIAKNHKTTVQRLVSLNSISNADVISIGDV
|   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |   |
5 ANHLSSTTVLSIGQTLITIPGSKSSSTSSSTTMKSGSSVYTVKSGDSLMLTANEFKNTVQRLKKNLGLS-SDLIRAGQK
      70      80      90      100     110     120     130

543      573      603      633      663      693      723      753
LKID-----NSTASQARAKSQPTTENSMSNNTGSSDSAAKEIAS*IKKVVILHRMDNIMEDINCLNLT*MATYILAKI
||:   :||:   :|   :|   |||   |||   |:   :   |:   :   :
10 LKVGSTVSSSSSSSKKSNKSSSSSKSSSNKSSSSSTGTGTVKLGDSLANKIANKVMSTARLKVNLNLEKSDITVYN
      150     160     170     180     190     200     210

```

SEQ ID 8712 (GBS166) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 30 (lane 2; MW 13.1kDa).

The GBS166-His fusion product was purified (Figure 200, lane 10) and used to immunise mice. The resulting antiserum was used for FACS (Figure 315), which confirmed that the protein is immunoaccessible on GBS bacteria.

SEQ ID 4 (GBS15) was expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 9 (lane 5; MW 44.8kDa), Figure 63 (lane 5; MW 44.8kDa) and Figure 66 (lane 7; MW 45kDa). It was also expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 10 (lane 4; MW 22.3kDa). It was also expressed as GBS15L, with SDS-PAGE analysis of total cell extract is shown in Figure 185 (lane 1; MW 50kDa).

Purified GBS15-GST is shown in Figure 91A, Figure 190 (lane 9), Figure 210 (lane 4) and Figure 245 (lanes 4 & 5).

The purified GBS15-GST fusion product was used to immunise mice (lane 1 + 2 products; 20µg/mouse). The resulting antiserum was used for Western blot (Figure 91B), FACS (Figure 91C), and in the *in vivo* passive protection assay (Table III). These tests confirm that the protein is immunoaccessible on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 3

A DNA sequence (GBSx0091) was identified in *S.galactiae* <SEQ ID 303> which encodes the amino acid sequence <SEQ ID 304>. Analysis of this protein sequence reveals the following:

```

Possible site: 32
>>> Seems to have no N-terminal signal sequence
INTEGRAL    Likelihood = -9.66    Transmembrane    22 - 38 ( 15 - 41)
----- Final Results -----
bacterial membrane --- Certainty=0.4864(Affirmative) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:CAA72096 GB:Y11213 hypothetical protein [Streptococcus thermophilus]
Identities = 149/274 (54%), Positives = 208/274 (75%), Gaps = 9/274 (3%)

Query: 23 FLVSLLSFGIFSLITPKSN--KLTKKDFLTNKKVILPNYVALGDSLTEGVGDTTSQGGF 80

```


- F + LL GI IIP S+ K+ K KK + YVA+GDSLT+GVGD++QGGF
 Sbjct: 5 FLLFLLLPVGLILFIIPSSQSSKSDIKRISVKKK-KVITYVAIGDSLTCGVGDSSNQGGF 63
- 5 Query: 81 VPLISESLHNRYSQVTSVNYGVSGNTSQQLIKRMITDPQIEKDLKADLLTLTVGGNDV 140
 VP+LS++L + ++QVT NYG+GNYS QILKRM I+DL+KA L+TLTVGGNDV
 Sbjct: 64 VPVLSQALSDFPNQVTPRNYGIAGNTSNQILKRMQKKDKIKRDLKAKIMTLTVGGNDV 123
- Query: 141 LAVIRKELSHLSIANSPEKPAEAYKRIKSLAKARQDNPKLPITYVLGIYNPFYLPFQIT 200
 + VI+ ++L+H+F K A Y++RL++I+ AR++N LPIY+GIIYNPFYLPF++T
 10 Sbjct: 124 IIVIKDNITNLMVNTFSKAVDYQKRLRQIILARKINKNTLPITYIIGIYNPFYLPFPIET 183
- Query: 201 KMQTVIDNNKATKEVVDASENVYFVPINDRLYKGINGKKGITSS-----SNSQASITN 254
 +MQT++DNVN++T+EV +NVYFVP+ND LYKGINGK G+T S + S N
 15 Sbjct: 184 EMQTVIDNNRSTREVSREYDNVYFVPINDRLYKGINGKGGVTSDEPISQPTKSSQDSIN 243
- Query: 255 DALFTGDHFHPNNIGYQIMSNAVMKINETRNK 286
 DALF DIIHFNN GYQIMS+A++++IN+T+K W
 Sbjct: 244 DALFERDHFHPNNIGYQIMSDAILKRINQTKEW 277
- 20 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 305> which encodes the amino acid sequence <SEQ ID 306>. Analysis of this protein sequence reveals the following:
- Possible site: 39
- 25 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -12.05 Transmembrane 18 - 34 (10 - 37)
- Final Results -----
 bacterial membrane --- Certainty=0.5919 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 30 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
- A related sequence was also identified in GAS <SEQ ID 9123> which encodes the amino acid sequence <SEQ ID 9124>. Analysis of this protein sequence reveals the following:
- Possible site: 33
- 35 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -12.05 Transmembrane 12 - 28
- Final Results -----
 bacterial membrane --- Certainty=0.5919 (Affirmative) < succ>
 40 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
- An alignment of the GAS and GBS proteins is shown below:
- Identities = 178/282 (63%), Positives = 218/282 (77%)
- 45 Query: 5 LLLNFMVNKKKILTGSLFPLVSLLSFGI FSLIIPKSNPKLTKKDPTKKVIPLAYVALG 64
 L LNFVMN + + +G FF++SL L+F + ++IIPKSN +L K DFL K+ + + YVA+G
 Sbjct: 1 LRLNFMVNNRHLFSGIFFVVISLCLAFLLNIIIPKSNRLKKSDFLKQVAILQYVALG 60
- 50 Query: 65 DSLTEGVGDITSGGGFVPLLSLHNRYSQVTSVNYGVSGNTSQQLIKRMITDPQIERK 124
 DSLTEGVGD T QGGFVPL+ L + V NYGVSG+TSQQIL RM QI+
 Sbjct: 61 DSLTEGVGDLTHQGGFVPLITNDLSEYFKANNVHONYGVSGDTSQQILDRMIQLS 120
- Query: 125 LKADLLTLTVGGNDVLAVIRKELSHLSIANSPEKPAEAYKRIKSLAKARQDNPKLPITY 184
 L+KAD++TLTVGGNDV+AVIRK L+ L ++SF KEA Y++RL++I+ AR+DN LPI+
 55 Sbjct: 121 LKADIMTLTVGGNDVMAVIRKDLQVSSFKKPAQYQKRLAQIILARKINKDLPIF 180
- Query: 185 VLGIYNPFYLPFQITMQQVIDNNKATKEVVDASENVYFVPINDRLYKGINGKKGITR 244
 +LGIYNPFYLPF++L+ MQ VID+NN TKEVV + VYFVPIND LYKGING+HGI
 60 Sbjct: 181 ILGIYNPFYLPFRLDMQKVIDDNRWTKTKEVGEYRNVYFVPINDRLYKGINGKQGVH 240
- Query: 245 SNSQASITNDALFTGDHFHPNNIGYQIMSNAVMKINETRNK 286
 SS Q +I NDALFTGDHFHPNN GYQIMSNAVMKIN + K

Sbjct: 241 SSGDQTTIVNDALFTGDHPFNNTGYQIMSNVMEKIKKHEK 282

A related GBS gene <SEQ ID 5> and protein <SEQ ID 6> were also identified. Analysis of this protein sequence reveals the following:

```

5  Lipop: Possible site: -1  Crend: 4
   SRCFLG: 0
   MoG: Length of UR: 24
       Peak Value of UR: 3.02
       Net Charge of CR: 3
10  MoG: Discrim Score: 12.27
   GvH: Signal Score (-7.5): -3.44
       Possible site: 22
   >>> Seems to have an uncleavable N-term signal seq
   Amino Acid Composition: calculated from 1
15  ALGM program count: 1 value: -9.66 threshold: 0.0
       INTEGRAL Likelihood = -9.66 Transmembrane 12 - 28 ( 5 - 31)
       PERIPHERAL Likelihood = 1.96 118
       modified ALGM score: 2.43
   icml HYPID: 7 CFF: 0.486
20
   *** Reasoning Step: 3

   ----- Final Results -----
       bacterial membrane --- Certainty=0.4864(Affirmative) < succ>
25       bacterial outside --- Certainty=0.0000(Not Clear) < succ>
       bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the databases:

```

30  56.0/80.3% over 272aa
   GP|1850894| hypothetical protein Insert characterized
   ORF02006 (367 - 1164 of 1467)
   GP|1850894|emb|CAA72096.1||Y11213 (5 - 277 of 280) hypothetical protein {Streptococcus
   thermophilus}
35  %Match = 30.8
   %Identity = 56.0 %Similarity = 80.2
   Matches = 150 Mismatches = 49 Conservative Sub.s = 65

141 171 201 231 261 291 321 351
40  AV*RPBANG I I L L K V P K H E K L K L A S P T V V K L I W L I T L E K N * L F * V L L Y P F * K L A Q S S K I L A V R M H L L A P M N K K I L

381 411 435 465 495 525 555 585
45  T G L S F F L V L L S F G I S P L S I I P K S N - - P K L T G C D F L T K K V I E L N V A L G D S L T S G V G D T S Q G G F V L L S E S L N R Y S Y Q
   : : : : | | : : | | : : | | : : | | : : | | : : | | : : | | : : | | : : | | : : | | : : | | : : | | : : | | : : | | : : | | : : | |
   S F A G F F L L L F V G I L I F I P S S H Q S S I G S D K I R S V K K - E K V T V A L G D S L T G V G D S S N G G F V P V L S Q A L S D F N W Q
       10 20 30 40 50 60 70

615 645 675 705 735 765 795 825
50  V T S N Y G V S G N T S Q Q I L K R M T T D P Q I E K D L E K A D L L T L T V G C N D V I A V I R K H S E L S L N S P E K D A P A Y K E R L K S I L A K A R
   : : | : : | | | : : | | | | : : | | | : : | | | : : | | | : : | | | : : | | | : : | | | : : | | | : : | | | : : | | | : : | | | : : | | | : : | | |
   V T P R N Y G I A G I Y N P F I N P F M T E Q T I V D N N R S T E E V S K E Y D N V F V F V N D L Y K G I N K G G V S D S Q P T K S S
       90 100 110 120 130 140 150

855 885 915 945 975 1005 1044
55  Q D N P K L P I Y V L G I Y N P F I N P F Q L T K M Q T V I D N N K A T K E V D A S E N V Y F V P I N D R L Y K G I N K E G I T - - - - - E S S N S
   : : | | | : : | | | | | | : : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : | : |
   K E N K T L P I Y I G I Y N P F I N P F M T E Q T I V D N N R S T E E V S K E Y D N V F V F V N D L Y K G I N K G G V S D S Q P T K S S
       170 180 190 200 210 220 230

1074 1104 1134 1164 1194 1224 1254 1284
60  Q A S I T I N D A L P T G D H P F N N I G Y Q I M S N A V M E K I N S T R K N P * F K F L E M G I S L I V C N * P F L H S D C K S L N S T * A * Y R K N F
   | : | | | | | | | | | | | | | | : : : : | : | : |
   Q D S L - N D A L F P E E D H P F N N I G Y Q I M S D A I L K R I N Q T K K W S G E
       250 260 270 280

```

65

SEQ ID 6 (GBS103) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 36 (lane 4; MW 32kDa).

The GBS103-His fusion product was purified (Figure 107A; see also Figure 201, lane 9) and used to immunise mice (lane 2+3 product; 18.5µg/mouse). The resulting antiserum was used for Western blot (Figure 107B), FACS (Figure 107C) and in the *in vivo* passive protection assay (Table III). These tests confirm that the protein is immunoaccessible on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

10 Example 4

A DNA sequence (GBSx1316) was identified in *S.agalactiae* <SEQ ID 3837> which encodes the amino acid sequence <SEQ ID 3838>. Analysis of this protein sequence reveals the following:

```

Possible site: 23
>>> Seems to have no N-terminal signal sequence
15   INTEGRAL    Likelihood = -4.30    Transmembrane 1058 -1074 (1056 -1075)

----- Final Results -----
           bacterial membrane --- Certainty=0.2720 (Affirmative) < succ>
           bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
20           bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

A related GBS gene <SEQ ID 7> and protein <SEQ ID 8> were also identified. Analysis of this protein sequence reveals the following:

```

Lipop: Possible site: -1    Crend: 10
MoG: Discrim Score:      -13.26
GVH: Signal Score (-7.5): -5.76
Possible site: 41
30   >>> Seems to have no N-terminal signal sequence
ALOM program   count: 1 value: -4.30 threshold: 0.0
           INTEGRAL    Likelihood = -4.30    Transmembrane 489 - 505 ( 487 - 506)
           PERIPHERAL Likelihood =  3.71      97
           modified ALOM score:  1.36

35   *** Reasoning Step: 3

----- Final Results -----
           bacterial membrane --- Certainty=0.2720 (Affirmative) < succ>
           bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
40           bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

LEXTG motif: 478-482

```

45 SEQ ID 8 (GBS195) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 24 (lane 8). It was also expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 31 (lane 5).

GBS195C was expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 175 (lane 6 & 7; MW 81kDa).

GBS195L was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 83 (lane 2; MW 123kDa).

GBS195LN was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 83 (lane 3; MW 66kDa).

- 5 GBS195-GST was purified as shown in Figure 198, lane 5. GBS195-His was purified as shown in Figure 222, lane 4-5. GBS195N-His was purified as shown in Figure 222, lane 6-7.

The GBS195-GST fusion product was purified (Figure 87A) and used to immunise mice (lane 1 product; 13.6µg/mouse). The resulting antiserum was used for Western blot (Figure 87B), FACS, and in the *in vivo* passive protection assay (Table III). These tests confirm that the protein is immunoaccessible on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 5

A DNA sequence (GBSx0002) was identified in *S.agalactiae* <SEQ ID 4043> which encodes the amino acid sequence <SEQ ID 4044>. This protein is predicted to be lipoprotein MlsA. Analysis of this protein sequence reveals the following:

```

Possible site: 19
>>> Seems to have no N-terminal signal sequence
----- Final Results -----
      bacterial cytoplasm --- Certainty=0.3361(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 9403> which encodes amino acid sequence <SEQ ID 9404> was also identified.

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 3177> which encodes the amino acid sequence <SEQ ID 3178>. Analysis of this protein sequence reveals the following:

```

Possible site: 13
>>> Seems to have no N-terminal signal sequence
----- Final Results -----
      bacterial cytoplasm --- Certainty=0.2412(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

```

Identities = 146/168 (86%), Positives = 161/168 (94%)

Query: 1  MNLENGIITYSKNTIAQLAKDPKNKATYENKRDYAVAKLEKLDKRAKSKFNAIPANKILI 60
      +NLENGIITYSKNTIAQLAKDPKNK TYENK AYVAKLEKLDKRAKSKF+AI NKILI
Sbjct: 107  LALENGIITYSKNTIAQLAKDPKNKITYENKLAAYVAKLEKLDKRAKSKFDIAENKILI 166

Query: 61  VTSEGCFKYPSKAYGVPSAYIWEINTEEGTDPQITSLVKLKKQVRPSALPVSSVDIKRP 120
      VTSEGCFKYPSKAYGVPSAYIWEINTEEGTDPQI+SL++KLK ++PSALPVSSVD+RP
Sbjct: 167  VTSEGCFKYPSKAYGVPSAYIWEINTEEGTDPQITSSLIETKLKVLKPSALPVSSVDIKRP 226

```

Query: 121 MKSVSRRESGIPTIAETFDLSIAKKGKGSYANMKWNLDKIEGLAK 168
 M+VS+SGIPIY+EIFDLSIAKKG+ GDSYANMKWNLDKIEGLAK
 Sbjct: 227 METVSKDGIPIYSEIFDLSIAKKGKGSYANMKWNLDKIEGLAK 274

- 5 SEQ ID 9404 (GBS679) was expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 164 (lane 7-9; MW 36kDa) and in Figure 188 (lane 8; MW 36kDa). Purified protein is shown in Figure 242, lanes 9 & 10.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

10 Example 6

A DNA sequence (GBSx0003) was identified in *S. agalactiae* <SEQ ID 8485> which encodes the amino acid sequence <SEQ ID 8486>. This protein is predicted to be ATP-binding protein MtsB. Analysis of this protein sequence reveals the following:

- Possible site: 55
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.2097(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 8765> which encodes the amino acid sequence <SEQ ID 8766>. Analysis of this protein sequence reveals the following:

- Possible site: 29
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1929(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

- Identities = 143/238 (60%), Positives = 186/238 (78%), Gaps = 2/238 (0%)
 Query: 1 MIISKHLGVSYDNNL-VLEDINLRLEGGSIIGILGPNAGKSTTANKALLGLVDSTGESGI 59
 MI + L V+YD N LE IN+ +EG I+GI+GPNAGKST MKA+L L+D G +
 Sbjct: 10 MITINNLGVYDGNALRINVTIGPPIVGIIGPNAGKSTVMKALINLDYQSHVTV 69
 Query: 60 GG-DLLELWLRVAYVQKTNIDYQFPITVGRCVSLGLYKERGLFKRLSKTDWEKVSRLVID 118
 G D L VAYVQ+ IDY FPITV ECV+LG Y + GLF+R + K +E+V +V+
 Sbjct: 70 DKGIDGRKLGHVAYVQKSNIDYNFPITVKECVLGVYSLGLFPRVGRKQFQVQDVKVLK 129
 Query: 119 QVGLRGFNRPINLALSGGQFQRMILMARCLVQEAUYIFLDEPFVGINDSBQIIVNLLKKL 178
 QVGL F +RPI +LGGGQFQRMILARCI+QD+DYIFLDEPFVGINDS+SB+IIV+LLK+L
 Sbjct: 130 QVGLDEPGHRPIKSLGGGQFQRMILARCI+QSDYIFLDEPFVGINDSBQIIVNLLKKL 189
 Query: 179 SKAGKLLVHHHLSKVHYPDQVQVILNRLHILACGSPIDQAFTRRNLSAAYGAILLQ 236
 AGK IL+VHIEDLSKV+HYFD++ILN+HL+A G + + FT + LS AYG+ +LG+
 Sbjct: 190 RMAGKTLIVHIEDLSKVHYPDKMLILNKLHVAIGNVCVFTVDTLSKAYGNHILGK 247

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 7

A DNA sequence (GBSx0004) was identified in *Sagalactiae* <SEQ ID 9> which encodes the amino acid sequence <SEQ ID 10>. Analysis of this protein sequence reveals the following:

Possible site: 28

>>> Seems to have an uncleavable N-term signal seq

----- Final Results -----

bacterial membrane	---	Certainty=0.0000 (Not Clear)	< succ>
bacterial outside	---	Certainty=0.0000 (Not Clear)	< succ>
bacterial cytoplasm	---	Certainty=0.0000 (Not Clear)	< succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 8

A DNA sequence (GBSx0005) was identified in *Sagalactiae* <SEQ ID 11> which encodes the amino acid sequence <SEQ ID 12>. This protein is predicted to be integral membrane protein MtsC (znuB). Analysis of this protein sequence reveals the following:

Lipop: Possible site: -1 Crend: 6

McG: Discrim Score: 3.77

GvH: Signal Score (-7.5): -0.47

Possible site: 45

>>> Seems to have a cleavable N-term signal seq.

INTEGRAL	Likelihood = -10.83	Transmembrane	138 - 154 (134 - 162)
INTEGRAL	Likelihood = -7.96	Transmembrane	60 - 76 (50 - 86)
INTEGRAL	Likelihood = -6.95	Transmembrane	95 - 111 (93 - 118)
INTEGRAL	Likelihood = -5.79	Transmembrane	180 - 196 (174 - 216)
INTEGRAL	Likelihood = -4.35	Transmembrane	198 - 214 (197 - 216)
INTEGRAL	Likelihood = -4.30	Transmembrane	250 - 266 (246 - 268)
INTEGRAL	Likelihood = -3.93	Transmembrane	222 - 238 (221 - 241)
PERIPHERAL	Likelihood = 5.94	116	
modified ALOM score:	2.67		

*** Reasoning Step: 3

----- Final Results -----

bacterial membrane	---	Certainty=0.5331 (Affirmative)	< succ>
bacterial outside	---	Certainty=0.0000 (Not Clear)	< succ>
bacterial cytoplasm	---	Certainty=0.0000 (Not Clear)	< succ>

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 13> which encodes the amino acid sequence <SEQ ID 14>. Analysis of this protein sequence reveals the following:

Possible site: 45

>>> Seems to have a cleavable N-term signal seq.

INTEGRAL	Likelihood = -11.25	Transmembrane	138 - 154 (134 - 163)
INTEGRAL	Likelihood = -9.08	Transmembrane	66 - 82 (50 - 86)
INTEGRAL	Likelihood = -6.79	Transmembrane	95 - 111 (93 - 118)
INTEGRAL	Likelihood = -5.63	Transmembrane	180 - 196 (176 - 216)
INTEGRAL	Likelihood = -4.73	Transmembrane	221 - 237 (218 - 241)
INTEGRAL	Likelihood = -4.35	Transmembrane	250 - 266 (246 - 268)
INTEGRAL	Likelihood = -4.35	Transmembrane	198 - 214 (197 - 216)
INTEGRAL	Likelihood = -2.81	Transmembrane	48 - 64 (47 - 64)

----- Final Results -----

bacterial membrane --- Certainty=0.5501 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

5 An alignment of the GAS and GBS proteins is shown below:

Identities = 224/275 (81%), Positives = 255/275 (92%)

```

Query: 1  MPTKFFEGLLTYHFLQNAITAIVIGIVAGVGCFTILRSMSLMKDAISHAVLQGVATSF 60
      M  KFFEGSL++YHFLQNA ITA+VIGIV+GAVGCFILRSMSLMKDAISHAVLQGVA+SF
10  Sbjct: 1  MSMKFFEGSLSYHFLQNALITAVVIGIVSGAVGCFITILRSMSLMKDAISHAVLQGVATSF 60

Query: 61  ILGINFFIGAIIVFGLLSSIIITYIKENSVIKGTATGITFSSFLAIGILLIGLANSTDL 120
      ILG+NFFIGAI+FGLL+S+IITYIKENSVIKGTATGITFSSFLAIG+ILIG+ANG+IDL
15  Sbjct: 61  ILGVNFFIGAIIFGLLASVIITYIKENSVIKGTATGITFSSFLAIGVILIGVANSSTDL 120

Query: 121  PHILFGNVLAVQDSKRYTIIVGLIVLITITFFKELLITSFDPVLAKSMGMVSVFHYL 180
      PHILFGNVLAVQDSK++TI V + VL +I++FFKELLITSFDP+LAKSMG++V+ YHYL
20  Sbjct: 121  PHILFGNVLAVQDSKMITIGVSIPIVVISLFFKELLITSFDPILAKSMGVKVNAYHYL 180

Query: 241  GYTFNIAAGSSIVLSTSTPMFLAFLSPKQSLPKK 275
      GYTFN+AAQSSIVLTS MFL++F SPKQ K+
25  Sbjct: 241  GYTFNVAAGSSIVLTSAMMFLISFFVSPKQGYLKR 275
  
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

30 Example 9

A DNA sequence (GBSx0006) was identified in *S.galactiae* <SEQ ID 15> which encodes the amino acid sequence <SEQ ID 16>. Analysis of this protein sequence reveals the following:

Possible site: 38

```

>>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.1280 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
40  Sbjct: 241  GYTFNVAAGSSIVLTSAMMFLISFFVSPKQGYLKR 275
  
```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 10

A DNA sequence (GBSx0007) was identified in *S.galactiae* <SEQ ID 17> which encodes the amino acid sequence <SEQ ID 18>. This protein is predicted to be peptidyl-prolyl cis-trans isomerase 10 (rotamase). Analysis of this protein sequence reveals the following:

```

50  Lipop Possible site: 19  Cread: 2
      McG: Discrim Score: 5.27
      GVH: Signal Score (-7.5): -4.14
      Possible site: 19
      >>> May be a lipoprotein
  
```

-51-

ALOM program count: 0 value: 9.34 threshold: 0.0
 PERIPHERAL Likelihood = 9.34 89
 modified ALOM score: -2.37

5 *** Reasoning Step: 3

----- Final Results -----

10 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAA19257 GB:AL023704 putative Cyclophilin-type peptidyl-prolyl
 cis-trans isomerase protein [Schizosaccharomyces pombe]
 15 Identities = 88/224 (39%), Positives = 123/224 (54%), Gaps = 46/224 (20%)
 Query: 50 NKKTQKALADKKAFFQLDKAVAKNEAQ-----VLKTSKGDINIKLFPKYAPL 98
 N TK D +D+ + + V NE + +I T++GDI+IKL+P+ AP
 Sbjct: 419 NMSTKFTLSDRDVINEQVLPVTNNEGRQENGNILLGKAAIHHTTQGDISTIKLYPEAPK 477
 20 Query: 99 AVENFLTHAKBTYANGLSFHRVVKDFMIQSGDENGDTGGKSIWNSKDKKDGNGQFVNE 158
 AV+NF THR+ GYY+ FHR+IK+FMIQ GDP GDGTGG+SIW KKD F +E
 Sbjct: 478 AVQNFTTHAENGYYDNTIFHRILIKNFMIQGGDFLGDGTGGESIW----KKD----PDE 528
 25 Query: 159 ISPLYLNIHG-SLAMAGAGDTNGSQFFITNGSQDHSKQLSDKKVPKVIKAYSBGQNS 217
 ISP L + R +++MAN+G +TNGSQFFIT P
 Sbjct: 529 ISPNLKHDRPPTVSMANSGPNTNGSQFFITTDL-----TFW 564
 30 Query: 218 LDGGYTVFQGVQISQMETVVKIASVEVTKSDQPKKIKTISIKVI 261
 LDG +T+F + +G++V +I E K D+P E I +I +
 Sbjct: 565 LDGKHTIFARAYAGLDVVHRIEQGETDKYDRPLEPTKIDILIV 608

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 19> which encodes the amino acid sequence <SEQ ID 20>. Analysis of this protein sequence reveals the following:

35 Possible site: 19

>>> May be a lipoprotein

----- Final Results -----

40 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

45 >GP:CA88542 GB:AL353818 putative protein (Arabidopsis thaliana)
 Identities = 83/186 (44%), Positives = 104/186 (55%), Gaps = 34/186 (18%)
 Query: 78 VVMRTSQGDITLKLFPKYAPLAVENFLTHAKGYDNLTPHRVINDFMQSGDQKGDGTG 137
 V+M T+ GDI +KL+D+ P VENE TH + GYDN FHRVI FMIQ+GDE GDGTG
 50 Sbjct: 476 VMHTTLDTHMKLYPEECPTVENFTTHCRNGYYDNLHLPHRVIRGFMQGTGDFLGDGTG 535
 Query: 138 GESIWKGDPKKQKNGCPVNEISPLYHIRG-ALAMAGAGNTNGSQFYINQKKNQSG 196
 G+SIW G P +E L H R L+MANAG NTNGSQF+I
 55 Sbjct: 536 GQSIW-----GREPDEPHKSLRHRDPPTLSMANAGPNTNGSQFFITT----- 578
 Query: 197 LSSNTYKPLIISAYEKGQNSLDGGYTVFQGVINDQNDVVDKIANTSINQNKPSQDITIT 256
 P ID +TVRG+V+ GNDVV I ++ND+P QD+I
 Sbjct: 579 -----VATPWLNDKHTVPGRVKGMVDVQSGIEKVKTDKDRPYQGVKIL 622
 60 Query: 257 SIDIVK 262
 ++ + K
 Sbjct: 623 NTVVPK 628

An alignment of the GAS and GBS proteins is shown below:

Identities = 172/267 (64%), Positives = 221/267 (82%)

```

5  Query: 1  MKKIIVLGLACVSIITLGGCESTIERSLKGDYVVDQKLAENSSKEATBQLNKKTKQALKAD 60
      1  MKK++ L L +S+L LS CBS++R++KGD+Y+D+K A+ S+ A++ + ++ALKAD
      Sbjct: 1  MKKILSLSLVAISLNLNLSACBSVDRAIKGDXYIDKTKAKESRAASKAYEESIQALKAD 60

10 Query: 61  KKAFFQLDKAVAKNERAQLIKTSKGDINIKLFFKYAPLAVENFLTHAKGYNYGLSPHEV 120
      61  FFQL K V K EA+V+++TS+GDI +KLFFKYAPLAVENFLTHAK+GY+ L+PHEV
      Sbjct: 61  ASQFFQLTKEVGKEEAKVVMRTSQGDIITLKLFFKYAPLAVENFLTHAKGYNYDNLTPHEV 120

15 Query: 121  IKDFMIQSGDPFMGSGTGKSIWNSKDKKDSGNFVNEISPYLYNIRGSLAMANAGADTN 180
      121  I DFMIQSGDP GDGTGG+SIW KD KKD+GNFVNEISPY+LY+IRG+LAMANAAGA+TN
      Sbjct: 121  INDPMIQSGDPKSGDTGGSINWRSKDPKDAKGNFVNEISPYLYHIRGALAMANAGANTN 180

20 Query: 181  GSQFFINQSQDHSKQLSDKKVPKVIKAYSBSGNPSLGGYTVFGQVISMETVDKIAS 240
      181  GSQF+INQ+++ SK LS PK II AY GNPSLGGYTVFGQVI GM+ VDKIA+
      Sbjct: 181  GSQFYINQNKIKQSGKSLSTNYPKPIISAYEBSGNPSLGGYTVFGQVIDGMDVVDKIAA 240

25 Query: 241  VEVTKSDQPKKEKIIITSIKVINKDYKFK 267
      241  + ++D+P++ ITITSI ++KDY+FK
      Sbjct: 241  TSINQNDKPEQDITITSIDIVKDYRFK 267

```

SEQ ID 18 (GBS205) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 51 (lane 13; MW 31kDa).

GBS205-His was purified as shown in Figure 206, lane 8.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 11

A DNA sequence (GBSx0008) was identified in *S.galactiae* <SEQ ID 21> which encodes the amino acid sequence <SEQ ID 22>. This protein is predicted to be sporulation protein SpoIII_E (ftsK). Analysis of this protein sequence reveals the following:

```

Lipop Possible site: -1  Crend: 10
MoG: Discrim Score: -22.83
GVH: Signal Score (-7.5): -7.13
Possible site: 39
>>> Seems to have no N-terminal signal sequence
ALOM program count: 5 value: -9.24 threshold: 0.0
40  INTEGRAL Likelihood = -9.24 Transmembrane 36 - 52 ( 27 - 60)
    INTEGRAL Likelihood = -9.18 Transmembrane 162 - 178 ( 154 - 188)
    INTEGRAL Likelihood = -4.04 Transmembrane 597 - 613 ( 595 - 615)
    INTEGRAL Likelihood = -3.77 Transmembrane 63 - 79 ( 58 - 83)
    INTEGRAL Likelihood = -2.60 Transmembrane 90 - 106 ( 88 - 108)
45  PERIPHERAL Likelihood = 1.32 136
    modified ALOM score: 2.35

*** Reasoning Step: 3

----- Final Results -----
50      bacterial membrane --- Certainty=0.4694 (Affirmative) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 10035> which encodes amino acid sequence <SEQ ID 10036> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

-53-

>GP:CB13553 GB:Z99112 DNA translocase [Bacillus subtilis]
Identities = 352/822 (42%), Positives = 508/822 (60%), Gaps = 70/822 (8%)

5 Query: 14 KTRRTKAEIRFORAIQRMITALVLITILFLPGIIRLGIFGIVTVNIRFMVGLSLAYLFIA 73
K + R + + + + Q I + + I + I I + LG+ G T + R F G L + L +
Sbjct: 3 KKKRKRKKQAKQKLNKYRIKGLCIAISIIAILQLGVVQQTIFYLRFFFAGWETILCLL 62

Query: 74 ATLTYLYFFKMLRKDSL- ---AGFLIASLGLLIEWHAYLFS ---MPILKDKIIRST 125
L + W + K SL+ AG +L+ H LF ++ +R+T
10 Sbjct: 63 GLLVLGVSLFWKKKTPSLTRRKAGLYCIATILLSHVQLFKMI/THKGSIEASVVRST 122

Query: 126 ARLIVSDLMQFKITVFAGGGMIGALIKYPIAFLFSNIGAYMIGVLFILGLFMSLEVV 185
L + D + + GGGM+GAL++ FLF++ G+ ++ ++ I+G+ L++ ++
15 Sbjct: 123 WELFLMVMXGSSASPDLGGMIGALLFAASHFLFASTGSGQIMAVMILIGMILVGRSLQ 182

Query: 186 DIVE-----PIR----AFKN--KVAEKHONKKEKFAKREMKALAEQERIEKAE 231
K + + + F I + AF + K + + Q + K + A + + K + + + + + E + +
15 Sbjct: 183 ETLKKVMPIGRFKEQWLAFLIDMKSFNSNMQSKKTKAPSKKKPARKKQMQPEPPD 242

Query: 232 EEAYLASVNPDPETGHELEDQAEDNLDLAPPEVSETSTPVFEP--ELIAYETSPONPLP 290
EE +V+ + I+ ++ N ++ P + + + PV +P + + ET Q + +
20 Sbjct: 243 EGGDYETVSLIHSEPIISSFSRNEBEE--SPVIEKRAEPVSKFLQDIQETGDC--ETVS 300

Query: 291 VEPTIYLEDYDSPINRKNREDEEMVYDLDLDDVDSIEDVDFTEKTIIVYKLPITDLFAP 350
P + E F I + + EN D Y++P++DL A
25 Sbjct: 301 APFMIFTE-----LENKD-----YEMPSLDLLAD 324

Query: 351 DKPNQSKKEDLVKNKIRVLESTFSPGIDVKVERAIEGSPVTKYIEKPAQVGRVNRIG 410
K Q + K + + N R LE TP+SFG+ KV + +GP+VTKYE+ P VGV+V++I N
30 Sbjct: 325 PKHTGQADKKNIYENARKLESTFQSGFGVAKVTVQVILGPAVTKYVYVFDGVKSVIYN 384

Query: 411 LSDDLALALAAKDVRISTPIPGKSLIGIEVNSRIATVSPRELWQES--DANPENLLEVP 469
LSDDLALALAAKD+RIE PIPGKS IGIEVNP+E+A VS +E + P + + L
35 Sbjct: 385 LSDDLALALAAKDRIEAPIPGKSAIGIEVNAEAVMSLKEVLSKINDRDPANVLIGL 444

Query: 470 GKAVNGNARNFNLARMPHLLVAGSTGSGKSVAVNGIISILMKARPQVKPMIDPDMVE 529
G+ ++G A L +MPLLAVG+TSGKSV VNGII+SIIM+A+P +VK NMIDPDMVE
40 Sbjct: 445 GRNISGEAVLAEALNMPHLLVAGATGSGKSVVNGIITSILMAKAPHEVNMIDPDMVE 504

Query: 530 LSVYNDIPHLLIPVVTNPKASKALQKVVDEMRNYELFSKIGVURNIAGVNTKVEFNAS 589
L+VYN IPHLL PVVT+P+KAS+AL+KVV+EME RYELFS G RNI GYN ++ N
40 Sbjct: 505 LNVYNGIPHLLAPVVTDPKASQALKKVVNEMRRYELFSHTGTRNIEGVNDYIKRANIE 564

Query: 590 SEBQKIPLEPLIVIVDELADLMVVASKEVDNAIRLQKARAAGIHILATQRPVSDVIS 649
KQ LP IVIVDELADLMVVAS +VED+I RL Q ARAAGIH+I+ATORPVSVDVIT+
45 Sbjct: 565 EGAKQPELPYIVIVDELADLMVVASVSDVDSITRLSQNARAAGIHILATQRPVSDVIT 624

Query: 650 GLIKANVPSRIAFVSSQDTSRTILDENGAELKLRGDMFLFKPIDENHPVRLQGSFISD 709
G+IKAN+PSRIAF+VSS TDSRTILD GAELKLRGDMFL P+ N PVR+QG+P+SD
50 Sbjct: 625 GVIKANIPSRIFAFVSSQDTSRTILDMGGAELKLRGDMFLPVGANKPVRVQGAFLSD 684

Query: 710 DVERIVGPIKDQAEADYDAFDPGEVSETDNGSGGGVPSDDPLFPEAKGLVLETKQAS 769
+VE++V + Q +A Y + P E +ET + +D L+EA L++ Q AS
55 Sbjct: 685 EYEVVVDHVITQQKAGVQSEMIPEETETES-----EVTDELYDEAVELVGMQAS 736

Query: 770 ASMIQRRLSVGNPNRATIMELEAAGVIGPABSTKPRKVLMT 811
SM+QRR +G+ RA RL++ +E GV+GP EG+KPR+VL++
Sbjct: 737 VSMIQRFRIGYTRAAKILDAMBERGVVGPYBGSKPREVLIS 778

46.5/66.5% over 775aa

OMNI|NT01B81964| sporulation protein SpoIIIE Insert characterized

ORF01349(340 - 2733 of 3048)
65 OMNI|NT01B81964(6 - 781 of 790) sporulation protein SpoIIIE
%Match = 29.6
%Identity = 46.4 %Similarity = 66.5
Matches = 352 Mismatches = 243 Conservative Sub.s = 152

90 120 150 180 210 240 270 300
TIN*LATT*S*YTDG*TKINNFHTYSLIKLLR*LYFIINF*IIYKSK**TYWGTC*NYDRIV*HELIEKVRNKYFT*N

330 360 390 420 450 480 510 540
MVFANKKKTKGGKKRPTKABIERQRAIQSMITALVLITLIFPGIIRLGIGPITVYNVRPMVGSLAYLPDAATLIYLY
| : | : : : | : | : : | : | : | : | : | : | : | : | : | :
VMSVAKKKRKRKRKKQAQAKINIKYEINGLCLTAISIALILGVGQQVQFYFLFRFPAGENFICLLSGLLVGV
10 20 30 40 50 60 70

10

[illegible]

15

786 816 846 894 924 954
KPIAFLFSNIGAYMGVLFIILGLFMSLSLEVYDIVE-----FIR---AF--KNKVAEKHQNKKKERFAKREMKKA
|||:::||:::||:::||::||::||::||::||::||::||::||::||::||::||::||::||::||::||
AAASHFLFASTGSQAIDAVYMILGMLVTRSLRQLTKLKWNSPIGRIPIKZWLAIFDDMKSFKSNMQSKCKTAKPSKQP

 170 180 190 200 210 220 230

2.5

[illegible]

30

```

1224      1254      281              1326      1356      1386      1416
DYDSPIPMRENDEENVIDLDD-DVDDSIENVDFTPT-----TLVYKLTIDLFAPDKPKNSKEKDLVRNIVLSE
      :   :   :   :   :   :   :   :   :   :   :   :   :   :   :   :   :   :   :   :
-----LDIQIPGTGDQTFVSAPNPTFLENKDYEMSLDLDLPKHTGQAADKNIKNIYENARKLR
      290              300              310              320              330              340

```

35

[illegible]

40

1683 1713 1743 1773 1803 1833 1863 1893
LMEQS-DANFENLLVPLGKAVNGNARSFNILARMPHLIVAGSTSGSKSVANGSISSILMKARDQVKFMIDPIOMVELS
| : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : :
VLESKLNDAPPDANVLGLGRNISGRAVLZLNKMPHLIVAGATSGSKSVANGIITSILMRAPKEVFQMIDPIOMVELN

440 450 460 470 480 490 500

45

[illegible]

50

[illegible]

2403 2433 2463 2493 2523 2553 2583 2613
PIDENHVRVLQGSFTSDDDVVERIVGFKIKQAPADYDAPFDGPGVSRITONGSGGGGVPSLDFEEAKGLVLTQKASAS
: | | | | | : | | | | | : | | | | | : | | | | | : | | | | | : | | | | | : | | | | |
PVGANKPVRGVGAFLSDSEYEVKVDHVITQKQAYCEHMPETTT-----HSKVTLYELDYDAVELLVGMGTASV
680 710 740 770 800 830 860 890

65

MIQRRLSVGNRATIRIMELEAAGVIGPAETKPKRVIMTPTPSS*EKTNLRNCRISPLCYEAMR*RRRLRGHETVI
|::||:::|::|:::|::|::|::|::|::|::: :
MLQRRFRIGYTAARLIDAMEERGVGVPYESSKPRFVLISKRYDRLSS

760 770 780 790

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 23> which encodes the amino acid sequence <SEQ ID 24>. Analysis of this protein sequence reveals the following:

5 Possible site: 51
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -9.45 Transmembrane 31 - 47 (25 - 55)
 INTEGRAL Likelihood = -7.17 Transmembrane 160 - 176 (153 - 183)
 INTEGRAL Likelihood = -4.99 Transmembrane 93 - 109 (86 - 111)
 10 INTEGRAL Likelihood = -4.04 Transmembrane 586 - 602 (584 - 604)
 INTEGRAL Likelihood = -1.22 Transmembrane 64 - 80 (64 - 80)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.4779(Affirmative) < succ>
 15 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the databases:

1GB:Z99112 DNA translocase [Bacillus subtilis] 601 e-170
 Identities = 354/816 (43%), Positives = 499/816 (60%), Gaps = 69/816 (8%)
 20 Query: 11 APKRLTKAEVEKQRAIKRMILSVLMALLLIFAMRLGVGVVTTYIMIRFLVGLSLAYPM 70
 A KKR ++ + KQ IK + +L + I A+L+LGV G T + RF G +
 Sbjct: 2 AKKKRKRKKQAKQINIKYELNSGLCIAISIIAIIQLGVVQQTFLYLFRRFAGEMFLICG 61
 25 Query: 71 FAWLIYLCPCWLRQKQKGM-----AGVVIAPIGLLVENHAFLEA-----MPRMQLQDIFLG 122
 L+ W ++ ++ AG+ +L+ H LF + +
 Sbjct: 62 LGLVLVGLSVLFWKKTKPSLLTRKAGLYCITASILLSHVQLFKNIHKGSISSAVVRN 121
 30 Query: 123 TARLITRDLALRVTFVGGGMI GALLYKPIAFLFSNIGSYFIFGLFILGLFIATPWDI 182
 T L D+ + +GGGM+GALL+ FLF++ GS + + IL+G+ L+T +
 Sbjct: 122 TWELFLMDMNGSSASPOLGGMGALLFAASHFLFASTGSGMAIVMILIGMILVTRSL 181
 35 Query: 183 YD-----VSHFVKEA---VDKLAVAYQENKEKRFIKREHRLQAKRALEKQAE 230
 + + P+KE +D+ +++ N+ K+ + +K A +KQ E
 Sbjct: 182 QETLKWMSPIGRFPIKEQMLAFIDDMK-SFKSNMQSS--KKTAPSKQKQARKKQCMSP 238
 40 Query: 231 EKRLAELTVPETGEIVDSQSQVSYDLAEDMT-KEPEILAYDSLKDDSTSLFQD--- 285
 E E G+ Y+ + EP I ++ +++E+ ++
 Sbjct: 239 EP-----PDEEGD-----YETVSLIHSEPIISSFDMSREERSFVIEKRAEP 281
 45 Query: 286 --EDLAYAHEEIGAYDSLALASSEDEMCMDPEVEVDFTPKTHLLYKLPITIDLEAPDKPK 343
 + L E G ++SA + E++ + Y++P+DL A K
 Sbjct: 282 VSKPLQDQIPETGQDQETVSPAPMTPIELEND-----YEMPDLGLADPKIT 328
 50 Query: 344 NQSEKNLVKRNKIKVLEDTQSFQIDVKVERKIGFSVTYKIEKPAVGVRVNRINSLADD 403
 Q +K + +N + LE TFQSGF+ KV + +GP+VTKYE+ P VGV+V+I NL+DD
 Sbjct: 329 GQADKKNIYEMARKERTFQSGFVKAKVTQVHLGPAVTYKYEYVDPGVKVKSVINLSD 388
 55 Query: 404 LALALAAKDVRIEAPIGKSLIGIEVPNSEIATVSPRELNEQS-DANPENLLEVPGLKAV 462
 LALALAAKD+RIEAPIGKSLIGIEVPN+E+A VS +E+ E + P+ + + LG+ +
 Sbjct: 389 LALALAAKDRIEAPIGKSLIGIEVPN+EAVMSLKEVLSKLNDRPDANVLIGLRNI 448
 60 Query: 463 NGNARSFNLRMPHLLVAGSTGSGKSVAVNGIISILMKARPDQVKNMIDPRNVLGSVY 522
 +G A L +MPHLLVAG+TSGSKSV VNGIIL+SIIL+A+P +VK NMIDPRNVL+VY
 Sbjct: 449 SGEAVLAEINMPHLLVAGTSGSGKSVVNGIITISILMRAPHEVKNMIDPRNVLNLY 508
 Query: 523 NDIPHLIPVVTNPRKASKALQKVVDENEMRYELFSKIGVNRNAGYNTKVEFNASSEQ 582
 N IPHLL PVVT+P+KAS+AL+KVV+EMS RYELFS G RNI GYN + + N K
 60 Sbjct: 509 NGIPHLLAVVTPDKKASQALKKVVNEMERREYELFSHTGTRNIEGVDNYIKRANSEGA 568
 Query: 583 QIPLPLIVVIVDELADLMMVASKEVEDAIRLGQKARAAGIHMLIATORPSVDVISGLIK 642
 Q LP IVVIVDELADLMMVAS +VED+I RL Q ARAAGIH+I+ATQRPVDVI+G+IK
 Sbjct: 569 QPELPYIVVIVDELADLMMVASSDVEDSITRLSQARAAGIHMLIATORPSVDVITGVIK 628

Query:	643 ANVPSRMFAVSSQDTSRTILDENGAEKLLRGDMLFKPIDENHPVRLQGSFISDDOVER AN+PSR+AF+VSS TDSRTILD GA EKLLRGDMLF P+ N PVR+QG+F+SIN+VH+
Sbjct:	629 ANIPSRIPAVSSQDTSRTILDMGAEKLLRGDMLFLPVGANKPVRVQCAPLSDDEVEK 688
5	Query: 703 IVNFIKDQTEADYDDAFDGPGEVSDNDGPGSGNGAAGBDFLPEERAKLVLETQKASAMI +V+ + Q +A Y + P R ++ + D L+ +EA L++ Q AS SM+
Sbjct:	689 VVDHVITQQAQYQEEIMPEETTHSEVT-----DELYDEAVELIVGMQYASVSM 740
10	Query: 763 QRELVSQGNENRTRIMDEERAGVIGPABTKPRRVL 798 QRR +G+ RA RL+D +KE GV+GP HG+KPR+VL
Sbjct:	741 QRRFRIGVTRAARLIDAMEERGUVGPGYEGSEKPRRVL 776
An alignment of the GAS and GBS proteins is shown below:	
15	Identities = 620/818 (75%), Positives = 701/818 (84%), Gaps = 25/818 (3%)
Query:	1 MVMFMANKKTKGKTKRPTKAEIERQRAIQRMITALVLTLILFGIIRLIGFVITVYNI MV +KK+ KK R TRAE+E+QRAI+RMI ++++ ++L F ++RLG+FG+T YN+I
Sbjct:	1 MVRKQRKKSAPKK--RLTKAEVKQKRAIKMILSVLMALLLIFAMLLGLGVGVVYTNNI 58
20	Query: 61 RFMVGSLAYLFIATLIYLYFFKWLKKSLVAGFLIASGLLIEWHAYLFSPMLIK 120 RF+VGS LAY F+ A LIYL+ FWLK+KD ++AG +LA LGLL+EWHA+LF+MP + D++
Sbjct:	59 RFLVGS LAYLFFMFAMILYLFCFKWLKQKQMGIAVGVIAPLGLLEWHAFLFAMPRMLDQD 118
25	Query: 121 ILRSTARLIVSDLNQFKITVFAAGGMLGALYKPIAFLEFSGNIGVYMLFIIIGLFIAS 180 I TARKL DL+ ++T F GGGMLGAL+YKPIAFLEFSGNIG+Y IG LFI+LGLFLIAS
Sbjct:	119 IFLGSTARLITRDLALRVTEFVGGGMLGALYKPIAFLEFSGNIGSYFIFGLFIIIGLFLAIT 178
30	Query: 181 SLEVDIVFIFAPKKNVAEGHFNKKERFAKREMKALBQERIERQKASEEYASLVN 240 ++YD+ F++ +K+A +++NK++RF KRE + AE+E +B+Q EEE LA+
Sbjct:	179 PWDIYDVSHFKEAVDKLAVAYQFNKKRPIKREHRLQAEKEALEKQAESEKRLAELT 238
35	Query: 241 VDPETGEILEDQADNLDLDPFEVSETSTFPVFEILAYETSPQNDPLV--EPTIYL 297 VDPETGEI+ED + ++E T EPEILAY++ ++D + E Y
Sbjct:	239 VDPETGEIVDSQEQ-----VSYDLAEDWTK--EPEILAYDSHLKODETSLFDQEDLAVA 291
40	Query: 298 ED---YDSPIPMRENDSEMYDLDLDDVDSDIENVDTPKTLTLYKLPITIDLFAEDIF 353 + YDS + + ++EM D+D V+ VDPETPT L+YKLPITIDLFAEDIF
Sbjct:	292 HEEIGAYDS-LSALASEDEN--DMDEPVE-----VDPTKTLHLLYKLPITIDLFAEDIF 342
45	Query: 354 KIQSKSEDLVRKNIHVLZETFRSGFDIVKVERAETGPGSVTKYEIKPAVGVRVNRISLSD 413 KIQSKSEK+LVRKNI+VLE+TP+SGFDIVKVERAETGPGSVTKYEIKPAVGVRVNRISLSD
Sbjct:	343 KIQSKSEDLVRKNIHVLZETPQSGFDIVKVERAETGPGSVTKYEIKPAVGVRVNRISLSD 402
50	Query: 414 DALALAAKDVRITETPIPKSLIGIEVPMSEIATVSPRELWQSDANPENLLEVPGLKAV 473 DALALAAKDVRIR PIPKSLIGIEVPMSEIATVSPRELWQSDANPENLLEVPGLKAV
Sbjct:	403 DALALAAKDVRITETPIPKSLIGIEVPMSEIATVSPRELWQSDANPENLLEVPGLKAV 462
55	Query: 474 NQNARSFNLRMPHLLVAGSTGSGKSAVANGIISILMKARPQVKFMIDPQWELSVY 533 NQNARSFNLRMPHLLVAGSTGSGKSAVANGIISILMKARPQVKFMIDPQWELSVY
Sbjct:	463 NQNARSFNLRMPHLLVAGSTGSGKSAVANGIISILMKARPQVKFMIDPQWELSVY 522
60	Query: 534 NDIPHILLIPVVTNPKASKALQKVVDENMRYELPSKIGVKNLAGYNTKVEENASSEQK 593 NDIPHILLIPVVTNPKASKALQKVVDENMRYELPSKIGVKNLAGYNTKVEENASSEQK
Sbjct:	523 NDIPHILLIPVVTNPKASKALQKVVDENMRYELPSKIGVKNLAGYNTKVEENASSEQK 562
65	Query: 594 QIPLPLIVVIVDELADLMVASKKEVEDAIIRLGQKARAAGIHMLIATQRFSDVVISGLIK 653 QIPLPLIVVIVDELADLMVASKKEVEDAIIRLGQKARAAGIHMLIATQRFSDVVISGLIK
Sbjct:	583 QIPLPLIVVIVDELADLMVASKKEVEDAIIRLGQKARAAGIHMLIATQRFSDVVISGLIK 642
70	Query: 654 ANVPSRIAPAVSSQDTSRTILDENGAEKLLRGDMLFKPIDENHPVRLQGSFISDDOVER 713 ANVPSR+APAVSSQDTSRTILDENGAEKLLRGDMLFKPIDENHPVRLQGSFISDDOVER
Sbjct:	643 ANVPSRMFAVSSQDTSRTILDENGAEKLLRGDMLFKPIDENHPVRLQGSFISDDOVER 702
75	Query: 714 IVNFIKDQTEADYDDAFDGPGEVSDNDGPGSGNGAAGBDFLPEERAKLVLETQKASAMI 773 IV FIKDQ EADYDDAFDGPGEVSDNDGPGSGNGAAGBDFLPEERAKLVLETQKASAMI
Sbjct:	703 IVNFIKDQTEADYDDAFDGPGEVSDNDGPGSGNGAAGBDFLPEERAKLVLETQKASAMI 762

Query: 774 QRRLSVGFNRATRLMRELSEAGVIGPAEGTKPRKVLMT 811
 QRRLSVGFNRATRLM+ELE AGVIGPAEGTKPRKVL T
 Sbjct: 763 QRRLSVGFNRATRLMRELSEAGVIGPAEGTKPRKVLQT 800

- 5 SEQ ID 22 (GBS272d) was expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 147 (lane 9; MW 55kDa + lane 10; MW 70kDa). It was also expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 147 (lane 11 & 13; MW 85kDa + lane 12; MW 74kDa).

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for
 10 vaccines or diagnostics.

Example 12

A DNA sequence (GBSx0009) was identified in *S. galactiae* <SEQ ID 25> which encodes the amino acid sequence <SEQ ID 26>. This protein is predicted to be para-aminobenzoate synthetase (pabB) (pabB). Analysis of this protein sequence reveals the following:

15 Possible site: 61
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 20 bacterial cytoplasm --- Certainty=0.4073 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

25 >GP:AAD07357 GS:AE000547 para-aminobenzoate synthetase (pabB)
 [Helicobacter pylori 26695]
 Identities = 204/580 (35%), Positives = 325/580 (55%), Gaps = 50/580 (8%)
 Query: 16 YRFKNPTKELIADTLRQLVLEVIKEVDYYQSNYYVVGYLSYEASAAF-DSHFVVSQCKLA 74
 30 +++ K+L A L + + + Y+V GYL YEA AF D +F+ L
 Sbjct: 6 FKYQKSVKKLTATNLNRLNALDIFISQNRNGYFV-GYILYEARLAFDLNRLNLSKNSYKVN 64
 Query: 75 GSHLAY---FTVHKDCNEAFPLSYENVLEADNNWTANVSQEQYQEAITANI KQIIRCGNTY 131
 E +++ E+ +P + +++ ++ Y + +K +++ G+TY
 35 Sbjct: 65 FBQFLERKKYSLEPLKEHAFYFKIH-----SSLDQKTYFKQFKAVERKLNKGDY 114
 Query: 132 QVNYTLELSQOLCSDPPFSYTERLMVBOGAGYNAYIDDKRILSVSPELFFKKK--DEVL 189
 QVN T++L + P V++ ++ Q + A+I + +LS SPELFF+ D +
 40 Sbjct: 115 QVNLTMDELDTKAKPKRVFVNVNNTPTPKAFINFGSVLSFSPELFFLEFLDTAI 174
 Query: 190 T--TRPWGTSARKPTQSDVARRDWLANDPKNRSNNMIVDLLRNNGRICDVGTVKVK 247
 T+PWGKT AR D R +L ND KNRSN+MIVDLLRND+ R+ +VKV
 45 Sbjct: 175 KIIITPKWGTIRKSNKLIDKKNRILFLQNDKNRSNNMIVDLLRNLRIALKNYSKVN 234
 Query: 248 KLCQVEQYATVWQNTSTIRGVLSPEVTIMSTIQALYPCGSITGAPKISTMAINELEKRP 307
 +L ++ +V+Q S IE L +L IF+AL+PCGS+TG PKI TM II LEKRP
 50 Sbjct: 235 QLFETISLSPVYQNISEIEAKLPLKTSLEIFKALPCGSVTGCPKIITMQIIESLEKRP 294
 Query: 308 RGIYGGITGLCMPDQAI PNVPRTVQMKGGQ--AYYVGGGITWESQTSYESTRQS 365
 RG+YOG IG+ + +A+P+VDIRT++ + + GVG G+T+S+ EYEE+ KS
 55 Sbjct: 295 RGVYCGAIGM-VEEKKALPSVPITRLKRVHRNFIHGVGSGVTYKAPKPKYESSFKS 353
 Query: 366 -AVLTRVNPKEQLITTRGV--TRNKILFSQQ--HVERLRSASYFAYSFDKSKFERELKK 420
 V+ ++ +P++ T ++ + KL + + H ERL+ S YF + +D++ + EL
 60 Sbjct: 354 FFWPMKI--EFEIVETNKKIKDKQKLEINNNNAHKRNMSTRYFNFYDENLDFEL-- 409
 Query: 421 YIHQLDEKDYR+KIMLDKTKGVTFPSVKQLVNLKKFPLTAESVVQDYPI-KLSPTTYFKTS 479
 EK+ L+++L+K GK+ E K L L + E+ + + PI K + F Y KT+

5
 10
 15
 20
 25
 30
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A related DNA sequence was identified in *S. pyogenes* <SEQ ID 27> which encodes the amino acid sequence <SEQ ID 28>. Analysis of this protein sequence reveals the following:

Possible site: 31

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2669 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 303/572 (52%), Positives = 406/572 (70%), Gaps = 1/572 (0%)

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Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 13

A DNA sequence (GBSx0010) was identified in *S. agalactiae* <SEQ ID 29> which encodes the amino acid sequence <SEQ ID 30>. Analysis of this protein sequence reveals the following:

```
Possible site: 20

>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.1564 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 31> which encodes the amino acid sequence <SEQ ID 32>. Analysis of this protein sequence reveals the following:

```
Possible site: 13

>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.5335 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

An alignment of the GAS and GBS proteins is shown below:

```
Identities = 220/267 (82%), Positives = 243/267 (90%)

Query: 10 LLLSITKTIARATYYQLKKLNKPNKDKAISKSDIQSIYDEHNGNYGYRRIYLELRNRGFI 69
      +LLEI ++R+TYYQ+K+L + +KD +K I+ IYDEH+GNYGYRRI++ELNRNGFV+
Sbjct: 1 MLEIILDLRSRTYYQVVKLAQGDGKIDELKHVIREIYDEHNGNYGYRRIHMLELRNRGFVV 60

Query: 70 NHKRVQGLAKSMGLTARIRRRKRYASYKGEVGKADNLIQRQFEGSKPYKCYTDTVEFA 129
      NHK+VQ LMK MGL ARIRRRKRY+SYKGEVGKADNLI+R FEGSKPYKCYTDTVE A
Sbjct: 61 NHKRVQRLMKVMGLAARIRRRKRYASYKGEVGKADNLIKRHFEGSKPYKCYTDTVELA 120

Query: 130 LPBGKLYLSPVLDGYNSEIIDFTLSRSPDLKQVQIMLEAFAAASSETILHSDQGQWYQ 189
      LPBGKLYLSPVLDGYNSEIIDFTLSRSP+LKQVQIMLE+ FFA SYS TILHSDQGQWYQ
Sbjct: 121 LPBGKLYLSPVLDGYNSEIIDFTLSRSPNLKQVQIMLEKTFPADSYSGTILHSDQGQWYQ 180

Query: 190 HKSYHQFLEDKGIRPSMRKGNPONGMESFFGLLKSEMFYGLEYSKSLDLEQAITD 249
      E+SYH FLE KGI SMRSKGNPONGMESFFGLLKSEMFYGLE +Y+SLD LE+AITD
Sbjct: 181 HQSYHDFLESKGIILASMRKGNPONGMESFFGLLKSEMFYGLLETTYQSLDLEQAITD 240

Query: 250 YIFPYNNKRIKAKIKGLSPVQYRTKSF 276
      YIFPYNNKRIKAKIKG SPVQYRTKSF
Sbjct: 241 YIFPYNNKRIKAKIKGSPVQYRTKSF 267
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 14

A DNA sequence (GBSx0011; GBSx2234) was identified in *S. agalactiae* <SEQ ID 33> which encodes the amino acid sequence <SEQ ID 34>. Analysis of this protein sequence reveals the following:

```
Possible site: 27
```


-60-

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

5 bacterial cytoplasm --- Certainty=0.3578 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 35> which encodes the amino acid sequence <SEQ ID 36>. Analysis of this protein sequence reveals the following:

10 Possible site: 25

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

15 bacterial cytoplasm --- Certainty=0.3869 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

20 Identities = 107/170 (62%), Positives = 134/170 (77%)

Query: 1 MGLSYEDKLEIYELRKIGMSNSQISQRYDVRISNLKYMIKLMERYGVEIKGRNEYYPF 60
 MK + E K++IYELR++G S IS++D+ S+LKYMI+L+DRYGV IV+K +N YY P
Sbjct: 1 MKPQETKVKIYELRQMGESIKSISKFDMAESDLKYMIRLIDRYGVTVIVQCKKHYYSP 60

25 Query: 61 ELKQEMIDKVLHGCSQLSVSLDYALNSCSITLNMWLSQFKKNGYTVIVEKTRGRPSIKGRK 120
 ELKQS+I+KVL I G SQ SLDYAL S+L+ W++Q+KKNGYTI+EK RGRPSIKGRK
Sbjct: 61 ELKQEIINKVLIDQSQKQTSLDYALFTSSMLSRWIAQYKKNGYTILEKPRGRPSIKGRK 120

30 Query: 121 RKKTEEMTELRLQENRERLRTENAFLLKRLDLRLDEALQSERQKOLE 170
 RKK EEMTE+ERLQ+ E R E NA LKKLR+ RLRDEA E+QK +
Sbjct: 121 RKNLEEMTEVERLQKLEYPRAENAVLKKLREYRLRDEAKLEQKQSPK 170

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 15

A DNA sequence (GBSx0012) was identified in *S. galactiae* <SEQ ID 37> which encodes the amino acid sequence <SEQ ID 38>. This protein is predicted to be oxyR protein. Analysis of this protein sequence reveals the following:

40 Possible site: 22

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

45 bacterial cytoplasm --- Certainty=0.1323 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 10033> which encodes amino acid sequence <SEQ ID 10034> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:CA91664 GB:Z67753 former trsE (rbcR homolog) [Odontella sinensis]
Identities = 72/259 (27%), Positives = 127/259 (48%), Gaps = 7/259 (2%)

55 Query: 5 QKIMYLESIELYSNITKAAHLFISQPYLSKVTKQLENELEIKLIQSQGQTPLITYVQGR 64
 Q+L L++I + T+AA LF+SQP LSK IK LE+ L I L+ + + LT AG+

Sbjct: 8 QQLRILKAIATEKSPTRAARVLFVSQPSLSQIKTLESRLNISLLNRNNVSLTQAGKL 67
 Query: 65 YLFYLKEIDMIEKMAKELYLRSDKKGRITLIGNSGLASSILANVLKFNLEHPEISVK 124
 +L Y + I + + + L +++ +G + +G + + + + VL F HP+ + + +
 5 Sbjct: 68 FLEYSERILALCESCRVINDLKTDRGNLIVGASQTIGTYLMRVLALFAQHNPQINTE 127
 Query: 125 LLENNQNISEQLVASGDIDLAV--GMAPILYKDGIASTTYRDELPMITPTSQLYNAEK 182
 + ++ + V GEID+AV G P + + DKL L+IP + +K
 10 Sbjct: 128 VHVDSRKIAKRVLEGDIDLAVVQGNIPPEIEKNLKVDFVNDZELLIIIPKSHFPALKKK 187
 Query: 183 RGQIIPFEYPISVLD-NEPLILTPLEYGIGKTTAQFYELHMSLNQMISTTVEPTAASLS 241
 + Y ++ + N + L I IA F + Q+ + + TA SL
 Sbjct: 188 KKINKDDLYHLNFITLANSNSTIRKIDNILLIQIA-PEPEQFNIIQMNSIEAKTPAVSL- 245
 15 Query: 242 LSGMGATFVPQTLHRYLD 260
 G+GA FV + I + ++
 Sbjct: 246 --GLGMAFVSSSAIEKEIE 262

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 39> which encodes the amino acid
 20 sequence <SEQ ID 40>. Analysis of this protein sequence reveals the following:

Possible site: 30
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -1.28 Transmembrane 109 - 125 (109 - 126)
 25 INTEGRAL Likelihood = -0.27 Transmembrane 146 - 162 (146 - 162)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.1510 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

30 The protein has homology with the following sequences in the databases:

>GP_AAC22434 GB:U32761 transcriptional regulator [Haemophilus influenzae Rd]
 Identities = 157/303 (51%), Positives = 221/303 (72%)
 35 Query: 2 IRQGBSYLDIKOIRYPIATVENHFMLSQAALLVVSQPTLSMMINDFEKRENVLFKRKR 61
 + +G +DI+ +RYF++IV+N FNLG+A++ LYVSOP LSMMI +PE REN++PKR
 Sbjct: 9 VLRGVMDIRHLRYFVSIVDNDNFMLSRASQNLVVSQPALSMMITFENRENTQIFKRAS 68
 40 Query: 62 GRIIGLTYLGDNYKDAQVLSYDYMFLKLDHSHKGLKGSINIGIPPLILSVFSEVMP 121
 G+IIGLT+ G+NYY+DA++V+ Y+DM L+ +G+I IGIPPL+LS VFS V+L
 Sbjct: 69 GKIIGLT+GAGNYRDAKEVIRKYNMRTNLYKSKDCKKGTITIGIPPLVLSRVFSSVLP 128
 45 Query: 122 KLILENPGIQFNVKEIGAYOLKNELLVGVSDVAVLSPGTIADNLVETYEIQRSELSVCL 181
 L+L+NP I F +KEIGAY LK+ELL+ VD+AVLL P I+ N++++ RI SEL++ L
 Sbjct: 129 HLILKNPDIINFIKEIGAYALSELLDKVDLAVLVPERISQNIIDGIEHSSSEALFL 188
 50 Query: 182 SPRHRLASKKVIQMEDLTDQQLALPDPSFVMEHLVLEACRHOVRNIIITSSSDFMNL 241
 SP+H LA K+ I W DL +++A+PD +FM+HH + EA ER+ P+I+L SS MDF+L+
 Sbjct: 189 SPKHVLAKKQQTWADLHQQMAIPDQTFMHHHLKEAFERNNCYPDVLSDSCMDPLLS 248
 55 Query: 242 STKINHNVLITCPKPTITELQKLDKICPMRPSISRWVLTIRAKKSYSEISYIMODLL 301
 + K N +LTI P P+ ELN K+ C +R P +V+L R R K Y+ +E YI D LL
 Sbjct: 249 AVKINKELWLTILEPMAKLYBSKEFLCRKIESPVFWKVLCLQRKQTVTHLEYEYIDKLL 308
 Query: 302 QSF 304
 ++F
 Sbjct: 309 EAF 311

An alignment of the GAS and GBS proteins is shown below:

60 Identities = 61/227 (26%), Positives = 111/227 (48%), Gaps = 10/227 (4%)
 Query: 9 YLEISRLYSNITKAANHLFTSPQYLSKVTKOLENLEIKLIQ-SQGHQVTLTYAGQRYLF 67
 ++ +E +N++AA L++SOP LS +I E +KL + +G LTY G Y
 Sbjct: 17 FTAIVENHFNLSQAALLVVSQPTLSMMINDFEKRENVLFKRKRGRIGLTYLGDNYK 76

Query: 68 YLKEIDMIERQNAKELVLRSDIKKGEITLGINSGHASSILANVLKPNLEHPEISVKLLE 127
 +++ + M +L+ KG I +GI + S + + V+PK I&+P I + E
 Sbjct: 77 DAQKVLSDYDMFLKLDHDSKGLKGSINIGIPPLILSVVPSEVMFKILLENPGIQPNVKE 136

Query: 128 NNONISEQLVASGDIDLAVGMAPILYKDGIAST-TIYRDELFLMIPITTSQLYNAEKGQOI 186
 + + G++D+AV ++P D + T I R EL + + +L A K + +
 Sbjct: 137 IGAYQLKNEILLVGNVDVAVLSPGTGIADNLVETIYEQISELCVCLSPRHRL--ASKK--V 192

Query: 187 IPFEYPISVLNEPLILTLPTSYGIGKTIQAQFYELHJMSLAQMTITST 233
 I +E L +E L L + + + + E H + N ++T+S+
 Sbjct: 193 IQWE----DLTDEQLALEDPSPFVHHLVLRACERHQVRPNITLSSS 235

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 16

A DNA sequence (GBSx0013) was identified in *S.galactiae* <SEQ ID 41> which encodes the amino acid sequence <SEQ ID 42>. This protein is predicted to be aminoacylase (cpsA). Analysis of this protein sequence reveals the following:

Possible site: 43

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.75 Transmembrane 385 - 401 (385 - 401)

----- Final Results -----
 bacterial membrane --- Certainty=0.1298(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAF36227 GB:A7168363 aminoacylase [Lactococcus lactis]
 Identities = 201/395 (50%), Positives = 274/395 (68%), Gaps = 5/395 (1%)

Query: 6 LRHQLFEKLDQKCDQMAIRRYLHENFELSFKETKTAAYISDFYKQKDCHVQTFQGGNG 65
 L + L L Q ++M+ IRR+LH+ FE+S+E +T YI FYK DC + G G
 Sbjct: 3 LNNLLNLTSLTQYENEMIQIRRLHQYFEISFQKETPKYIMGFYKELDCPKLIGKGF-G 61

Query: 66 VVVDIYGDKATDKPKIKHIALRADFDALPIQEBTGLSPASKTAGVMHACGHDAHTAYLLIL 125
 ++VDI G K+ K +ALRADFDAL I E+ LSP S GVMHACGHDAHTAYL++L
 Sbjct: 62 IIVDIGGKSG---KTLALRADFDALAFENDLSFKSVNPGVMHACGHDAHTAYLMLV 117

Query: 126 AESLIELKSEFSGHIRILHQPAAEVPGGAKAMTEAGCLDGLDAVLGIHVMSTMEEGTVQ 185
 A L++K E G +RI+HQPAAEVP GSGAKMI+AG LDG+D ++G+HVM++T+ G +
 Sbjct: 118 ARELVKIKQLPGRAVIVHQPAAEVSFGGAKMIKAGALGDVNMIGVHMTTIKTGVIA 177

Query: 186 YHAGPIQTGRATFKVILQGGKQGHGSMHRANDTIVAASSPVMAAQITVSRVNPFDATAV 245
 YH CTGR+ F + ++G GGH SMP +ND IVAAS FV QT++SRR++PPD V
 Sbjct: 178 YHNKTKCTGRSGNFITTIKSGNGHASMPLQNSDAIVAASYPVTELTQTVISRIDPFDMGTV 237

Query: 246 TIGSPDGGKGSANVIKDSVLEGDVRVMSSETRGVVEERFKRILOGIAQTVGVSYQLDYQN 305
 TIGSPDGS N I+D V L+GDVR+M E TR V+ ++ K+I G+ T+GV +DY +
 Sbjct: 238 TIGSPDGGSGFNAIQDKVLLKGDVRMMKETTTRKVRDQVQKLAGGVGVTEGEVIVDYD 297

Query: 306 DYPVLVNNSEVTQKAVNSLKSVAIKRILDOVIDCDEQTPSEDFAYYQITIPACFYVGAHE 365
 +YPLV N+ +T V +SLK I E+ +++D PQ PSEDF+YY Q +P+ FFY+GA
 Sbjct: 298 NYPVLNSENLTHTVVDLSKQNI SEVNNIVDLGEQNSEDFSYGQVVESTFFYIQAQP 357

Query: 366 EQQFYYPHRRHPKQIAESSLMVSAKSMATAALAML 400
 E YEPH P F++ E S+++AK++AT + L
 Sbjct: 358 EDGKNYPHHSPLFKMNEKSLIAAKAVATVINYL 392

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 17

- 5 A DNA sequence (GBSx0014) was identified in *S.galactiae* <SEQ ID 43> which encodes the amino acid sequence <SEQ ID 44>. This protein is predicted to be drug transporter. Analysis of this protein sequence reveals the following:

```

Lipop: Possible site: -1  Crend: 8
McG: Discrim Score:      6.19
GVH: Signal score (-7.5): -0.899999
    Possible site: 31
>>> Seems to have a cleavable N-term signal seq.
ALOM program count: 11 value: -12.15 threshold: 0.0
15  INTEGRAL Likelihood = -12.15 Transmembrane 169 - 185 ( 166 - 190)
    INTEGRAL Likelihood = -8.86 Transmembrane 229 - 245 ( 224 - 250)
    INTEGRAL Likelihood = -8.65 Transmembrane 82 - 98 ( 78 - 111)
    INTEGRAL Likelihood = -8.60 Transmembrane 436 - 452 ( 428 - 457)
    INTEGRAL Likelihood = -7.48 Transmembrane 202 - 218 ( 198 - 222)
20  INTEGRAL Likelihood = -4.99 Transmembrane 334 - 350 ( 332 - 352)
    INTEGRAL Likelihood = -4.88 Transmembrane 358 - 374 ( 354 - 376)
    INTEGRAL Likelihood = -4.09 Transmembrane 301 - 317 ( 301 - 317)
    INTEGRAL Likelihood = -2.81 Transmembrane 102 - 118 ( 101 - 119)
    INTEGRAL Likelihood = -2.71 Transmembrane 52 - 68 ( 50 - 70)
25  INTEGRAL Likelihood = -1.70 Transmembrane 271 - 287 ( 270 - 288)
    PERIPHERAL Likelihood = 0.32 401
modified ALOM score: 2.93

*** Reasoning Step: 3

30  ----- Final Results -----
        bacterial membrane --- Certainty=0.5861 (Affirmative) < succ>
        bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
        bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

35  The protein has homology with the following sequences in the GENPEPT database:

>GP:CAE02058 GB:Z79702 hypothetical protein Rv2333c [Mycobacterium tuberculosis]
Identities = 118/405 (29%), Positives = 199/405 (49%), Gaps = 9/405 (2%)

40  Query: 13 KLLVGIVLAVLSFWLPAQS-ILNMG-PDVQSSLSGISSGMDIGVSTALFSGLFIVTVGG 70
    +L L I + F + F + I+N+ PD+Q S + + V+S +L +F+I+
    Sbjct: 5 QLLTLIATGLGLFWIFLDALVINVALPDIQRSFAVGEDGLQVNVASYSLGAVFVMSAAT 64

Query: 71 LADRLGRVKFTFIGLCLNIIGSLILVLANGAVLFIMGRIFQGLAAMPIMPSTMALVKTTY 130
    LAD GR ++ IG+ L +GS+ LA + R OGL RA + +++ALV +
45  Sbjct: 65 LADLGRRRRWYLGIVSLPTLGSIAQLAFSLAVLTANGAQLGMAVSVTSLAVGSAAF 124

Query: 131 -DGKDRQRAVSPWSIGSWGSGSLCSYFGGAVASTLQWRVYFISF-IASVVSFLILGTP 188
    + K++ RA+ W+ + G+ GG + GWR +F ++ ++V FL +
    Sbjct: 125 PEAKRKARAIGTWIASIGTTGTPTLGLLVDQNGWRSIFVNLMLGALVLPFLCYVE 184

50  Query: 189 ESKNVGQKTHFDYLGILIIPIISMLSLNIGISMAQEHGLMNVIPLSLPTVMIGLGFVLYTV 248
    ES N + FD G ++FI+++ +L + + G +V + + +G LF ++
    Sbjct: 185 ESKN-ERARRFDLSGQLLFTIVAVGALVAVIEGPDIGWTSVQTIVMLMTAVGCALEFVL 243

55  Query: 249 ETRKNSPFIDHFNRPY-LGATISNPLKAVAGTLIVNTYMQGQRQLTPKVAGENSL 307
    E R SN +D LF + Y L + AV G L++ ++Q R TP V G M L
    Sbjct: 244 ERRSNPMDLTFPDTSYALAIATICTVFVAVYGMILLTQFLQVRSYTPSVTGIMIL 303

Query: 308 GYLAVCVLIAIRVGEKTLQRFGARKMLLGAMSTFVGIFLMTLVNIQPLEYLVLPVFGYAL 367
60  + V I + ++ R GAR P+L G +G+ ++ + LV VG L

```

Sbjct: 304 PFSAAVAIVSPLVGHIVGRIGARVPILAGLCMMUGLLMLIPSEHRS---ALVLVGLGL 360

Query: 368 FGTLGLIYATPSTDTAIISSIPNEKVGSAAGIYKMASSLGGAIGVA 412
G+G+ + TP T A++++P E+ G ASGI ++G IG A

Sbjct: 361 CGSGVALCLTPIITVMTAVPAERAGNAGSAGMSAQRAIGSTIGFA 405

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 45> which encodes the amino acid sequence <SEQ ID 46>. Analysis of this protein sequence reveals the following:

Possible site: 61

>>> Seems to have an uncleavable N-term signal seq

INTEGRAL	Likelihood = -8.28	Transmembrane	169 - 185 (165 - 189)
INTEGRAL	Likelihood = -8.23	Transmembrane	12 - 28 (11 - 32)
INTEGRAL	Likelihood = -8.17	Transmembrane	429 - 445 (423 - 450)
INTEGRAL	Likelihood = -6.64	Transmembrane	203 - 219 (200 - 222)
INTEGRAL	Likelihood = -5.41	Transmembrane	227 - 243 (225 - 245)
INTEGRAL	Likelihood = -3.72	Transmembrane	82 - 98 (80 - 99)
INTEGRAL	Likelihood = -3.72	Transmembrane	136 - 152 (135 - 155)
INTEGRAL	Likelihood = -2.92	Transmembrane	302 - 318 (299 - 319)
INTEGRAL	Likelihood = -2.55	Transmembrane	261 - 277 (261 - 277)
INTEGRAL	Likelihood = -2.07	Transmembrane	331 - 347 (331 - 347)
INTEGRAL	Likelihood = -1.06	Transmembrane	56 - 72 (56 - 72)
INTEGRAL	Likelihood = -0.96	Transmembrane	351 - 367 (351 - 368)
INTEGRAL	Likelihood = -0.37	Transmembrane	104 - 120 (103 - 120)

----- Final Results -----

bacterial membrane --- Certainty=0.4312 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

iGB:A7250422 ORFC [Oenococcus oeni] 271 1e-71
Identities = 152/445 (34%), Positives = 248/445 (55%), Gaps = 7/445 (1%)

Query: 1 MSHHQQTQVSKQTIMAIIAIALIQFSGILSETSMNVTFPTIMSVYQLPINSLQNMITYILL 60
M Q VS +AI+ +A + F G+L ETSNMVTFPTIM + + LN +QN+TT YLL
Sbjct: 1 MQKDNQPVSLHVKLAILGLAGLAFQGVLIETSMNVTFPTIMAQCFSLINKVQNLITYALL 60

Query: 61 AVAIMTTSATLKKQVREPLFPNATGLTFPGTILAVLTQSFAMLLARIPQGITGLVM 120
VA ++ +A ++K + +FF A LF G I + L + F I+L R+ Q + TGL +
Sbjct: 61 LVAATISIAAFIEKRFPIKKIPFWAGLLPIIGVCSALAPNELLITGLRLIQALSTGLAI 120

Query: 121 PQMFNIILERVGHKVLGFWPGFAGLIISLAPAFGPTYGGFMISHFSQWIFICILFVPLI 180
P + I+++P K G +M ++ P+ GPTYGG + SN+ IF +LP+ LI
Sbjct: 121 PLLITEIMQQIPQKKQGSYMLVENLLLQPSLGPTYGGVITQDLSENRLIPWFLPILGILI 180

Query: 181 AGILAYYYLDESFPVSEKVPFDWLPALFALSISLTSALLAITSLE-NGSVNMLYGLFILTFSF 239
A ++ ++E K+PF W FI+L ++L S +A+ + G ++ + G ++
Sbjct: 181 AWLIGLSFIEQKSSPKIPFANKQIPSLITLALLSITTVAVINNAGIYGTWSIKFYGLPIIAV 240

Query: 240 IL---FLYKNLTAKQPLDIRILKIPSLTFLGILPFPVQLINLGINFLTPNFVMEKIAN 296
IL F+ + ++Q + I I K L+ +F+ Q I L + PL PN+ +
Sbjct: 241 IILIVFIKLSITNSRQALLISISIPKKGWFCPLLIYPLIQFIQLSLTFLPNYAQLILKKG 300

Query: 297 SSQAGMVLPLPGTILGALLAPAGKLYDQKGRISLYLGNALFSLSLITMITLQTRHFMPLP 356
+G++LL G+L A+L P G++ + ++ L +G S I T+ R+ +
Sbjct: 301 VMISGIMLLCSLISAILQLTGRMLDSFSVKIPLVIGAFFLITSTISPTIFQRLVSFVL 360

Query: 357 FTLLYILFTFGENNGVNNISLATAIRELPARKKADATATPQMMQQA/GAIGTAMAS-LIAN 415
LY+++ G + PNNSL A++LP + +D A+P +QQ+AG+IGT++AS L+AN
Sbjct: 361 IAAIYVITYMIGFSPFVNNISUTYALQKLPKLTSGNAVFNTLQQYAGTSIGTSVASALLAN 420

Query: 416 SQAESFTSGVQSVYLL*TFIALLDFT 440
T G QS Y +L+FI

Sbjct: 421 GIG--TDGQSNYTGSRHIFILNFI 443

An alignment of the GAS and GBS proteins is shown below:

Identities = 91/369 (24%), Positives = 160/369 (42%), Gaps = 14/369 (3%)

```

5  Query: 82  FIGLCINIIIGSLLVLVLANGAVLPINCRIPQGLAAAPIMPSTMAUVKTYDGKDRQRAVSF 141
    F+ L G+L VL + ++ RIFQG+ +MP ++ + F
    Sbjct: 83  FMATGLTFGTTLAVLTQSFAMILLARIPQIGITGLVMPQMFNIIILERVMPKVLPMGF 142

10  Query: 142  WSGISWGSGGLCSYPOGAVASTLGRVVFIPSIILASVVSPLLLIGTPESKNGKQKTHIFY 201
    + +OG + S W+ ++FI + +++ +L E V +K FD+
    Sbjct: 143  AGLIISLAPARGPTTYGGPMISHPSWQITPICILPVPLIAGIIAYYYLEDSPSEKVPFDM 202

    Query: 202  LGLIIFIISMLSINIGISMAQEHGLMVIPLSLPTVMILGIVLPVYVETRKSNSIFDPL 261
    L I IS+ S + I+ + E+G +N+ L LF ++ P+LF Y F+D +
    Sbjct: 203  LAFIALSISLTSALLAIT-SLENGSVNLYILGLF---ILSFILFLYKNLTAKQFFLDIRI 258

    Query: 262  FENRFLGATISNPLNAV-AGILIVINTVMQQGRQLTPKVAGEMSL-GYLVCVLIAIRV 319
    + I F+ + G + ++ + AG + L G L+ L+A
    Sbjct: 259  LKIPSLTFGLIPFFVQQLNLGNLINFITPFIIVMEKIANSSQKQMVLLPOTLLGALLAPAF 318

20  Query: 320  GEKILQRPGARKPMLIGAMSTFVGIVPLMTLVNIQGPLYLVLPV-VGVALETQGLIYATP 378
    G K+ + GAR + LG + + +MTL Q +++L P + Y LF G +
    Sbjct: 319  G-KLYDQKQARLSNLYGNALPSLSLITMTL---QTRHFMILLPTLLYLPTFGRMGFNN 374

25  Query: 379  STDTAISSIPNEKVGSGSIYIMASSLGGALGVATSIATYHAFSGMADPHKAALGGLIN 438
    S TAI + P EK A+ I+ +M GR+G A + I ++ A+P +L
    Sbjct: 375  SLATAIRELPKKNADATAIPQMMQCFAGALGTAMASLIANS---QAEFTSGVGVILLEF 431

    Query: 439  LVFCSLSIL 447
    +F L +
    Sbjct: 432  TIFALLDFI 440
  
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

35 Example 18

A DNA sequence (GBSx0015) was identified in *S. galactiae* <SEQ ID 47> which encodes the amino acid sequence <SEQ ID 48>. This protein is predicted to be transposase. Analysis of this protein sequence reveals the following:

```

40  Possible site: 45
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
    bacterial cytoplasm --- Certainty=0.3116 (Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
    bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
  
```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S. pyogenes*.

50 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 19

A DNA sequence (GBSx0016) was identified in *S.agalactiae* <SEQ ID 49> which encodes the amino acid sequence <SEQ ID 50>. This protein is predicted to be L11 protein (rplK). Analysis of this protein sequence reveals the following:

```

5   Possible site: 21
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
10  bacterial cytoplasm --- Certainty=0.1859(Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
    bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

15  >GP:CAA53739 GB:X76134 L11 protein [Staphylococcus carnosus]
    Identities = 117/139 (84%), Positives = 129/139 (92%)

    Query: 1   MAKKVEKLVKLQIPAGKATPAPPVGFALGQAGINIMGPTKEFNARTADQAGMIIPVVISV 60
              MAKKVEK+VKLQIPAGKA PAPPVGFALGQAG+NIMGF KEFNART +QAG+IIPV ISV
20  Sbjct: 1   MAKKVEKLVKLQIPAGKANEAPPVGFALGQAGVNINGPCKEFNARTQEQAGLIIPVEISV 60

    Query: 61  YEDKSFDFITKTTPAAVLLKKAAGVEKSGSEPNKTKVATITRAQVQEIATKMPDLAAN 120
              YED+SF FITKTTPFA VLLKKAAGVEKSGSEPNK KVAT-T+ QV+EIA+TKMPDLNAA+
25  Sbjct: 61  YEDRSFTFITKTTPAAPVLLKKAAGVEKSGSEPNKNKVATVTKDQVREIAQTKMPDLNAAAD 120

    Query: 121 LESAMRMIEGTARSMGFTV 139
              E+AMR+IEGTAREMG TV
30  Sbjct: 121 EEAAMRIIEGTARSMGITV 139

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 51> which encodes the amino acid sequence <SEQ ID 52>. Analysis of this protein sequence reveals the following:

```

    Possible site: 45

    >>> Seems to have no N-terminal signal sequence

35  ----- Final Results -----
    bacterial cytoplasm --- Certainty=0.4276(Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
    bacterial outside --- Certainty=0.0000(Not Clear) < succ>
40

```

An alignment of the GAS and GBS proteins is shown below:

```

    Identities = 136/141 (96%), Positives = 139/141 (98%)

    Query: 1   MAKKVEKLVKLQIPAGKATPAPPVGFALGQAGINIMGPTKEFNARTADQAGMIIPVVISV 60
              MAKKVEKLVKLQIPAGKATPAPPVGFALGQAGINIMGPTKEFNARTADQAGMIIPVVISV
45  Sbjct: 25  MAKKVEKLVKLQIPAGKATPAPPVGFALGQAGINIMGPTKEFNARTADQAGMIIPVVISV 84

    Query: 61  YEDKSFDFITKTTPAAVLLKKAAGVEKSGSEPNKTKVATITRAQVQEIATKMPDLAAN 120
              YEDKSFDFITKTTPAAVLLKKAAGVEKSG EN TKVAT+IRAQVQEIATKMPDLAAN
50  Sbjct: 85  YEDKSFDFITKTTPAAVLLKKAAGVEKSGSEPNKTKVATVIRAQVQEIATKMPDLAAN 144

    Query: 121 LESAMRMIEGTARSMGFTVTD 141
              +E+AMRMIEGTARSMGFTVTD
55  Sbjct: 145 IEAAMRMIEGTARSMGFTVTD 165

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 20

A DNA sequence (GBSx0017) was identified in *S. galactiae* <SEQ ID 53> which encodes the amino acid sequence <SEQ ID 54>. This protein is predicted to be ribosomal protein L1 (rplA). Analysis of this protein sequence reveals the following:

```

5   Possible site: 30
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
10  bacterial cytoplasm --- Certainty=0.2285(Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
    bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

15  >GP:CAB11879 GBS:Z99104 ribosomal protein L1 (BL1) [Bacillus subtilis]
    Identities = 144/228 (63%), Positives = 177/228 (77%)

    Query: 1  MAKKSIGNLRRAALEKIDSTKAYSVEEAVALKETNFAKFDATVEVSYNLMDVKKADQQIR 60
    Sbjct: 1  MAKKGKCTVEAAKLVDHSAKADYSEAVALKETNFAKFDATVEVAFRLGVDPSSNHHQQIR 60

    Query: 61  GAVVLPAFTGKTSEVLVPARGAKAEEAKAGADFVGEEDLVAKIQGWLDFDVVIATPDM 120
    Sbjct: 61  GA+VLP GTGKT EVLVFA+G KA+EA+AGALFVG+ D + KIQ GW DEDV++ATPDM
    GAVVLPAFTGKTSEVLVPARGAKAEEAKAGADFVGEEDLVAKIQGWLDFDVVIATPDM 120

25  Query: 121  MALVGRIGRLVGLSPNLMNPNTGTIVMDVAKAVESKGGKITRYADKAGNVQALIGKVSF 180
    Sbjct: 121  M VQ++GRLG+ LMPNPKTGTIV +V KA+ E K GK+ YR DKAGN+ IGKVSF
    MGEVGIKIGLSPKGLMNPNTGTIVTVEKAIKEIKAGKVEIRVDKAGNTHVPIGKVSF 180

30  Query: 181  DDAKLVDNPKAFNDVIVKAPATAKGTIYTNLSITTTQGVGKIVDPNSL 228
    Sbjct: 181  D KLV+NF+ D I+KAKPA AKG Y+ N+++T+T G G+RVD ++
    EDEKLVDNPTTHYITLTKAPAAKGVYKQVAVTSTMSPGVKVESST 228

```

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 55> which encodes the amino acid sequence <SEQ ID 56>. Analysis of this protein sequence reveals the following:

```

35  Possible site: 22
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
40  bacterial cytoplasm --- Certainty=0.2309(Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
    bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

45 An alignment of the GAS and GBS proteins is shown below:

```

    Identities = 208/229 (90%), Positives = 220/229 (95%)

    Query: 1  MAKKSIGNLRRAALEKIDSTKAYSVEEAVALKETNFAKFDATVEVSYNLMDVKKADQQIR 60
    Sbjct: 1  MAKKSIGNLRRAALEKIDSTKAYSVEEAVALKETNFAKFDATVEVSYNLMDVKKADQQIR 60

    Query: 61  GAVVLPAFTGKTSEVLVPARGAKAEEAKAGADFVGEEDLVAKIQGWLDFDVVIATPDM 120
    Sbjct: 61  GAVVLPAFTGKTSEVLVPARGAKAEEAKAGADFVGEEDLVAKIQGWLDFDVVIATPDM 120

55  Query: 121  MALVGRIGRLVGLSPNLMNPNTGTIVMDVAKAVESKGGKITRYADKAGNVQALIGKVSF 180
    Sbjct: 121  MA+VGRIGRLVGLSPNLMNPNTGTIVMDVAKAVESKGGKITRYADKAGNVQALIGKVSF
    MALVGRIGRLVGLSPNLMNPNTGTIVMDVAKAVESKGGKITRYADKAGNVQALIGKVSF 180

60  Query: 181  DDAKLVDNPKAFNDVIVKAPATAKGTIYTNLSITTTQGVGKIVDPNSL 229
    Sbjct: 181  D KLV+NFKAP+DV+ KAKPATAKGTIY+ N+SIT+TQGVGKIVDPNSL

```


Sbjct: 181 DADKLVENPKAFHDMAKAKPATAKGTMYANVSITSTQGVGIKVPNSL 229

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

5 Example 21

A DNA sequence (GBSx0018) was identified in *S. galactiae* <SEQ ID 57> which encodes the amino acid sequence <SEQ ID 58>. Analysis of this protein sequence reveals the following:

Possible site: 25

10 >>> May be a lipoprotein

----- Final Results -----

15 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 10029> which encodes amino acid sequence <SEQ ID 10030> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

20 >GP:BAE04286 GB:AP001509 nickel transport system (nickel-binding
protein) [Bacillus halodurans]
Identities = 209/541 (38%), Positives = 324/541 (59%), Gaps = 14/541 (2%)

25 Query: 5 RRNILLSTCLMVLTACHSDS----KSHKNSDK-LTLAWGEDFGDVPNHPYNDQF 59
R+ ILL + L+ L C +S + N+K +T +W D G +NPH YNP Q
Sbjct: 6 RKLILLFVILSISSILVGCASESGTVSNEGEENTEKSITFSWPRDIGMPNHPVNSQL 65

Query: 60 VIQMVYEGLVRYGQNGKIEPALAKSNISIQGKTYTFKLENA-KYSDGSNFAANVKRN 118
Q M+YE LV Y + G+++P LA SW+IS+DGK YTFKLE ++SDG+ FNA VK+N
30 Sbjct: 66 FQSGMIYEPVSVYTGEBELQPHLADSWTISDGKEYTFKLEGVQPSDGTFFNIVKIN 125

Query: 119 FDSIPSKENRGHMFNLNQLNENYALNQSTFBEIKLQAYSATLYDLSMIRPIRFLSDS 178
FD+ S+ H+W + N LE +++ TF++ LK+ Y L DL+++RP+RFL ++
35 Sbjct: 126 FDTWIEHSSL--HSLGVMNVLEKTEVVEFTFQVLKEPYYPALQDLAVRPRVFLGEA 183

Query: 179 AFPKGGDDITKQVKKPIGTGQWVVKSKKQNEYITFKRNENYMGKKPKLKEVTVKVIPDAQ 238
FP DT++ +K+PIGTG W++ KQ+EY F RN NYWG+ PK+ +VTVK+IPDA+
Sbjct: 184 GFPPDGDTSQ-GIKEPIGTGVMMLSDYKQDIYAVFTRNPNYMGESPKIDKVTVKIIPDAE 242

40 Query: 239 TRALAFESGDVLIYNGHIGLITPAQYTKDKKYVTAISQPMSTRLLLNKAKESIFODKK 298
TR LAFESG++DL+G G+I +D F Q + +Y T +S+P+ TR LLLN D +
Sbjct: 243 TRVLA FESGELDLIFGSEVISMDAFNQLKESGQYVTLSEFVGTRELLNTSNEKLADLR 302

45 Query: 299 VRQAMNHAIKVSIAKNTPFROTEKPADTIFPSKSTSHSIAKNIPYSYNDKANQLLDQAGM 358
VR A+H +K +++ G+E+ AD I S + ++D + P Y+V++AN LD+AGM
Sbjct: 303 VRLALHHGPNKQWVVEGVTGLGLEEKADNLSSTNPFYTDLDVEPIEYDVQANAYDLRAGM 362

Query: 359 KMGDK-VREKDGKTLRLPYLATKATKDLVTFQGSWRKIGINVSILAMSHEDDYAN 417
++ K VREK-G+ L L L Y T K + Q EW IG+ + + +E
50 Sbjct: 363 ELPAKTVREKNGEQLLELTYDKTDLQKAMAFNQBAWAIGVLDITGLSELTIQIR 422

Query: 418 AKKGNFDMMLTYSGAPWDPHAWMSALTAKADHGHENPINALENLAKTMDRLIKSALVD 477
+G+FD+ Y++GAP+DPH++++A+A G E R A NL+ K E+D +++ L
Sbjct: 423 RRAGDFDVFQWNYGAPYDPHSFIN-VVREAGWGVAE--AHNLKSKKEELDEQVATLAS 479

55 Query: 478 PKRENVDRTYKKVLELLHDEAVVILPYTQSVISYVRKGDFTMRFAPEENSPFLRYIEKNN 538
R Y + L L +++V++P+Y VY+++ F + P I+ +N
Sbjct: 480 TDIETREGLYSILNTLQSGSVFVPISYIKCTVYVQE-NVNEFTFPAIRDEHPFNGIDVSN 539

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 59> which encodes the amino acid sequence <SEQ ID 60>. Analysis of this protein sequence reveals the following:

Possible site: 24

>>> May be a lipoprotein

----- Final Results -----

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 131/497 (26%), Positives = 220/497 (43%), Gaps = 55/497 (11%)

```

15 Query: 8 ILLSITCLLMVILTACHSQDSKSHKIN-----SDKLTLAWGEDFGDVPNPHRYNP-DQFVI 61
    I L +T L++V AC Q ++ + D+L ++ G PH ++P D++ +
Sbjct: 13 ITLFLTLGLILV---ACQQKPKQTKERQKCRPKDELIVSMGAKL---PHEFDPKDRYGV 65

20 Query: 62 QD--MVYEGLVRYGDKGKIEPALAKSWISQDGKTYTFKLKNA-KYSDGNSFNAALNVIC 117
    + + + L++ I+ LAK++ S+DG T++F L + K+S+G A +VK
Sbjct: 66 HNEGNIHTSHLLKRSPELDIKGELAKTYHLSDEGLTWSFDLHDDFKFSNGEPTADDVKF 125

Query: 118 NFDSIFSKNIRGNHNFNLTNQLENYRALNQSTFEIKLKQAYSATLYDLSMIRPIRFLSD 177
    +D + + + ++LT ++N + + + I L +A+S L+ I PI
Sbjct: 126 TYDML-----KADGKANDLTF-IKNVEVVGKQVNIHLTEASTPTAQLTEI-PI----- 173

Query: 178 SAFPKG--DDTKKNVKKPIQGTQGVVSKSKQNEYITFKRNNYWGKKPKLKEVTVKVIP 235
    PK +D K N PIG+G ++VK K E F R N + GKPK K+ T V+
Sbjct: 174 --VPKKHYNDKYKSN---PIGSGPYVKEYKAGEQAI FVENPYWHGKKPYFKKWT-WVLL 227

30 Query: 236 DAQTRALAFESGDVDLYNGIIGLDLFAQYTK----DKKYVTAISQPMSTRLLLLNAKE 291
    D T A ESGDVE+Y + D + T+ V +S P + ++ ++ +
Sbjct: 228 DENTALAALBESGDVMYATPELA-DKKVKGTRLLDIPSDNVRGLSLPYVKGKGVITDSD 286

35 Query: 292 -----SIFQDKVRQAMRHAIKVSIAQNTFRGTEKPADTIFSKTSKSHDAKLNPFYSN 345
    + D +R+A+ +++ + G KPA +I K T + K
Sbjct: 287 GYPVGNVDVTSDFAIRKALTIGLNKQKVLDTVLNGYKGKPAYSIIDK-TPFWNPKTAIKDKK 345

40 Query: 346 VDKANQLLDQAGWQMGKDKVREKDGKTLTLRLPYIATKATDKLVLYTFQGSWMRKIGINVS 405
    V KA QLL +AGWK D R+K L Y +L + + +G +
Sbjct: 346 VAKAKQLLTAKGWKQADGSRKIGDLDAAPDLYPYTINDQLRANLAVEVABQAKALGITIK 405

Query: 406 LIAMBEDDYANAKGNFDMMLTYSWGAPWDPHAMWALTAADHGHNPENIALENLATK 465
    L A W + D L Y+ G +S + A G NI N I
Sbjct: 406 LKASN---WDENATKSHDSALLYAGORHEAQQFYSEHHPSLAGKGV-TNITFYVNPVT 460

Query: 466 E-MRLIKSALVDPKKE 481
    + +D+ + S+ +D E
Sbjct: 461 KYLDKAMTSSDLDKANE 477

```

A related GBS gene <SEQ ID 8469> and protein <SEQ ID 8470> were also identified. Analysis of this protein sequence reveals the following:

```

55 Lipop: Possible site: 22 Crend: 5
    MoG: Discrim Score: 7.69
    GVH: Signal Score (-7.5): -3.34
    Possible site: 25
    >>> May be a lipoprotein
    ALOM program count: 0 value: 7.21 threshold: 0.0
    PERIPHERAL Likelihood = 7.21 273
60 modified ALOM score: -1.94

```

*** Reasoning Step: 3

[illegible]

There is also homology to SEQ ID 318. An alignment of the GAS and GBS sequences follows:

Identities = 44/186 (23%), Positives = 78/186 (41%), Gaps = 27/186 (14%)

```

15 Query: 65 VQDMV-DGLLENDYGNLVPBLAKDWCSKDGIAITYTTVLRDSWSVYTADGEYAPVTAE 123
    VI MV +GL+ + G + P+AK W +S+DG TTY+ LR+ +DG + +
    Sbjct: 57 VIQDVFVIGLVRVYGDGKIPALAKGMSIQDGKTYTFKLNA--KYSDGNSFNANVK 113

Query: 124 DFTVTLGHAVDKSDALYVVEDSIQNLKAYQNGEVDKFGVGVKALDDQVYTLNKPFSY 183
    + + + + +N +AL+ T + L + +
20 Sbjct: 114 RNDSDIPKSNRGNHNVNLTQLN-----YRALNQSTPEKLK--QAY 156

Query: 184 WNSKITTSVLFPPVAKFLKS---KKGDPGTTDPSSILVNGAYPLGAPTSKSSMEFHKNE 239
    S T Y + +PL KQ D + + G + + + + F +NE
Sbjct: 157 --SATLYDLSMIRFIRFLGSDAPFKGDITTKKNVKKPICTGGVGVSKKQNEVITPKNE 214

25 Query: 240 NYWDAK 245
    NYW K
    Sbjct: 215 NYMGKK 220

```

30 SEQ ID 8470 (GBS186) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 35 (lane 7; MW 60kDa). It was also expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 41 (lane 6; MW 85.7kDa).

GBS186-GST was purified as shown in Figure 202, lane 4.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for
35 vaccines or diagnostics.

Example 22

A DNA sequence (GBSx0019) was identified in *S.agalactiae* <SEQ ID 61> which encodes the amino acid sequence <SEQ ID 62>. Analysis of this protein sequence reveals the following:

```

40      Possible site: 37

>>> Seems to have a cleavable N-term signal seq.
INTEGRAL    Likelihood = -5.95    Transmembrane    101 - 117 ( 99 - 123)
INTEGRAL    Likelihood = -4.73    Transmembrane    276 - 292 ( 275 - 293)
45    INTEGRAL    Likelihood = -1.12    Transmembrane    232 - 248 ( 232 - 248)
INTEGRAL    Likelihood = -0.96    Transmembrane    151 - 167 ( 150 - 169)

----- Final Results -----
          bacterial membrane --- Certainty=0.3378(Affirmative) < succ>
          bacterial outside --- Certainty=0.0000(Not Clear) < succ>
50    bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

55  >GF:BA04287 GB:AF001509 nickel transport system (permease)
    [Bacillus halodurans]
    Identities = 119/304 (39%), Positives = 174/304 (57%)

Query: 5   SSIKKKLAFALFFSLILFTLILKLSVNSAENYLRSLKISVSPEALKEARHYLGDK 64

```

S I K+I + + F + F+ I+LS V+ AE YL + I + E L E H GLD+
 5 Subjct: 3 SYIAKRIFAVIDIVLFAIFIMFVIRLSFVDPDAEAYLTAANHPTTELLAEKHHKSFGLDQ 62
 Query: 65 PLWKQYWLWFOKALTGDFGYSYVLRPLVDLVLQRFIATLFLGTSAFLIVITISTPLGW 124
 P+ QY K DFG+SYV FV D V R ATL L S+ L V I S PLG
 Subjct: 63 PMAVQYVQTIVKVFOLDFGHEVYINQFVWDEVIARMDATLQAVSSIFLAVLISPLGFL 122
 Query: 125 AGLHESARSDELIRFLSPSSVSMFNWVAYLLMLLFSAKINLLPVSGNDLQSLILPSIT 184
 + +++++ D R L\$+ S+P FW+ YLL+ ES KINL PV G L+LP+T
 10 Subjct: 123 SAIYKNSLIDRPSRLLSYLGASIPQFWLGYLLFPFVSVKINLFPVBRGSGWAHVLVITVT 182
 Query: 185 LSPSTVGQYIALIRKALISQENRSLNVENARLGRVKERYIVTHHLRNALPAINTALSLTW 244
 LS + + Y L+R + + + + V AR RG+KE+ I+ H+I+ A+ + + T L +
 15 Subjct: 183 LSLALIAIYTRILRASVLEQMOESYVLVARTGIRKEKVINVKHVLKLAISPVITSLGMIV 242
 Query: 245 VYLLTGSIIIVEIFSWNGIGRLFVTSIRTSDLFVIQACMLIFOTLFLANNFMTQCFMNV 304
 LLTG+IIV\$++FSW G GR FV ++ D+PVIQ +L+ LP+ N + +
 Subjct: 243 GKLLTGTIIVEQVFSWPGFRYFVDALFNRDIFVIQCTVLLAACLFIQCNLIVDIVLQAM 302
 20 Query: 305 DPRL 308
 DPR+
 Subjct: 303 DPRI 305

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 63> which encodes the amino acid
 25 sequence <SEQ ID 64>. Analysis of this protein sequence reveals the following:

Possible site: 40
 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -7.27 Transmembrane 290 - 306 (287 - 313)
 30 INTEGRAL Likelihood = -6.37 Transmembrane 12 - 28 (4 - 33)
 INTEGRAL Likelihood = -5.09 Transmembrane 105 - 121 (100 - 128)
 INTEGRAL Likelihood = -5.26 Transmembrane 145 - 161 (142 - 172)
 INTEGRAL Likelihood = -2.39 Transmembrane 191 - 207 (190 - 208)
 ----- Final Results -----
 35 bacterial membrane --- Certainty=0.3909(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

40 Identities = 102/324 (31%), Positives = 167/324 (51%), Gaps = 28/324 (8%)
 Query: 7 IIKKILSAFLALFPFISLLTFILIKLSTVN---SAENYLRLSKISVSPRALKEAHYGLD 63
 II KI+ +P +S+LTF+L+K S V+ ++ NY S++P K H+ GLD
 Subjct: 8 IIWKIIRCVTLIFGVSGLTVFLVKQSPVDPVWASVNY---DTSILTPAQYKAIAHHGLD 63
 45 Query: 64 KPLWKQYWLWFOKALTGDFGYSYVLRPLVDLVLQRFIATLFLGTSAFLIVITISTPLGW 123
 KP QY++W + + GD G S V R FV D++ R A+ L +++L I LG
 Subjct: 64 KPALVQYFNLNQLGQDLGTSILVYRQVSDIIRSRAGASFLIMGLSWILSGILGFLGT 123
 50 Query: 124 WAGLHESARSDELIRFLSPSSVSMFNWVAYLLMLLFSAKINLLPVSGNDL----- 175
 + H+ D ++R+ S+ +S+P FW+ + +L+FS +L P+ + +
 Subjct: 124 LSAFHQGLKLDVRVMFVSYLIQSVPTFWIGLIFLILFVSGVLSSEYVLFARAGETQWQIFKHCLR 183
 55 Query: 176 -----QSLILPSITLSPSTVGQYIALIRKALISQENRSLNVENARLGRVKERYIVTHHLR 230
 + L+LP TLS + R + S V AR RG + I HH LR
 Subjct: 184 LADRVKHNMLPVPFLLSILGIANVTLFTRTKMSVLSSEYVLFARAGETQWQIFKHCLR 243
 Query: 231 NALPAINTALSLTWVY---LLTGSIIIVEIFSWNGIGRLFVTSIRTSDLFVIQACMLIG 287
 N AI+ A++L + Y L GS++ E+FS+ G+G + SD P++ A ++I G
 60 Subjct: 244 N---ATVPATTLHPSYGRFLGSGVLAEQVFSYPLGSLTLEAGLKSITPLLAIAVMI-G 299
 Query: 288 TLF+L-ANNFMTQCFMNVDPRLRK 310
 TLF+ A N + + +P+LR+
 Subjct: 300 TLFVAGNLADIINSINPQLRR 323

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 23

A DNA sequence (GBSx0020) was identified in *S.galactiae* <SEQ ID 65> which encodes the amino acid sequence <SEQ ID 66>. This protein is predicted to be nickel transport system (permease). Analysis of this protein sequence reveals the following:

Possible site: 14

```
>>> Seems to have a cleavable N-term signal seq.
10  INTEGRAL    Likelihood = -7.64    Transmembrane    57 - 73 ( 51 - 80)
    INTEGRAL    Likelihood = -6.85    Transmembrane    173 - 169 ( 169 - 194)
    INTEGRAL    Likelihood = -5.79    Transmembrane    94 - 110 ( 86 - 112)
    INTEGRAL    Likelihood = -1.44    Transmembrane    221 - 237 ( 221 - 238)
15  INTEGRAL    Likelihood = -1.33    Transmembrane    118 - 134 ( 118 - 134)

----- Final Results -----
        bacterial membrane --- Certainty=0.4057(Affirmative) < succ>
        bacterial outside --- Certainty=0.0000(Not Clear) < succ>
20  bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>
```

The protein has homology with the following sequences in the GENPEPT database:

```
>GP:BAB04288 GB:AF001509 nickel transport system (permease)
[Bacillus halodurans]
25  Identities = 103/239 (43%), Positives = 157/239 (65%)

Query: 6  AIFAPILSSFDQYVLSQKLLAPNVHLLSTDLQGRDVLSELLYGARYSLFLAIIISLL 65
      AI AP ++ DP V+L- KLL P+ + LGTDQLCR LSRLL+GAR SL A +I +
Sbjct: 29  AILAPWIAHPDFIQVNLAKLLPPSWYFLGTDQLCRNLSELLFGARVSLGFLAIFIS 88

30  Query: 66  ELTIGMFVGLIVGWYQKLELFLWIANIILAPPSFLSLATVIGLHGLGNLIFAIVFV 125
      L IG+ VG I G+ G +++ + ++AFP+ +L L VG+ G GL ++ A+V V
Sbjct: 89  SLGITLLVGAIGYRGWIDSVLMRFCEGVNAPFNILVVLGLVGLPGGLMQVVLALVIV 148

35  Query: 126  EWVYYAKLMINIAKSAKPEYVINAQIMGLSVWHILRKHIFFPVYOFILVMVLMNIGNII 185
      +WVYYA++ +++ S K++ ++ A+I G S W I+R+HI P V FI+V+ + +G I
Sbjct: 149  QMVYYARNFRSMIVSLKEQNFITAARISGSSFWKILRRHIIPNVLFPIVVGITLEMGNWAI 208

Query: 186  LMISGFSFLGIGVQPNVTEWGMMLHDARGYFKTATWMLSPGIAIFLVPFSFNTLGDAI 244
      + IS SFLG+G+QF EWG M+H+ + + R+ +ML FGI I L V +FN LG++
40  Sbjct: 209  MDISALSLGLGIQFFTEWGMIMHEGKSFRISHPMLLPGIMILLVVMVTFNVLGSL 267
```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 67> which encodes the amino acid sequence <SEQ ID 68>. Analysis of this protein sequence reveals the following:

```
Possible site: 39
45  >>> Seems to have an unclesavable N-term signal seq
    INTEGRAL    Likelihood = -7.80    Transmembrane    182 - 198 ( 180 - 204)
    INTEGRAL    Likelihood = -7.38    Transmembrane    77 - 93 ( 69 - 98)
    INTEGRAL    Likelihood = -7.06    Transmembrane    112 - 128 ( 104 - 132)
    INTEGRAL    Likelihood = -6.16    Transmembrane    8 - 24 ( 7 - 31)
50  INTEGRAL    Likelihood = -5.10    Transmembrane    239 - 255 ( 235 - 258)

----- Final Results -----
        bacterial membrane --- Certainty=0.4121(Affirmative) < succ>
        bacterial outside --- Certainty=0.0000(Not Clear) < succ>
55  bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>
```

An alignment of the GAS and GBS proteins is shown below:

Identities = 61/246 (24%), Positives = 127/246 (50%), Gaps = 1/246 (0%)

-74-

Query: 2 LVISAIFAPILSSFPQYVDLSQKILAPNNVHLIGTDQLGRDVLRLLYGARYSLFLAII 61
 L+S+ + + P + + + LAP+ HL GTD LGRD+ R+G +SL+ ++
 5 Sbjct: 19 LILSILAINLYFPYKPTLEINAAIKNLAPSINHLPGTDLGLRDMFVKTIKGLYPSLQVGLL 78

Query: 62 ISLLELTGMFVGLIVGWQCKENLFLWIANIILAPPSFLSLATVGLIHLGLGNLIFA 121
 +L+ + + G++G ++ + W+ ++ + P+ + + ++G G+ +I A
 Sbjct: 79 GALMGVFLATVFGVLGHLGNSLIDKILAMLVDLFIGMPLHIFMILISFVVGKGAQGVIIA 138

Query: 122 IVFVENVYAKLMINLVKSAKKEPVVINAQIMGLSVNHLIKHIFPFVYQPIIWMVMI 181
 W A+L+ N V K+ +V ++ MG + +I+R HI P+ I + ++
 10 Sbjct: 139 TAVTHWPSLARLIRNEVYDLNKAIFVQLSKSMGKTPYIIVRHILPLASQIFIGFILF 198

Query: 182 GNIILMISGFSFLGIGVQPNVTEWGMHLHDARGYFRTAT+MMMLSPGLAIFLTVFSPNTL 240
 ++IL + +FLG G+ G+L+A + W+++ PG+ L+V +P+T+
 15 Sbjct: 199 PHVILHEASMTFLGPGLSAEQPSVGIIISRAKHSISLGNWLVIFPDLVLLVNAFDTI 258

Query: 241 GDAIDK 246
 G+++ K
 20 Sbjct: 259 GESLKK 264

A related GBS gene <SEQ ID 8473> and protein <SEQ ID 8474> were also identified. Analysis of this protein sequence reveals the following:

25 Lipop: Possible site: -1 Crend: 0
 MOG: Discrim Score: 7.56
 GVH: Signal Score (-7.5): -1.15
 Possible site: 14
 >>> Seems to have a cleavable N-term signal seq.
 ALOM program count: 5 value: -7.64 threshold: 0.0
 30 INTEGRAL Likelihood = -7.64 Transmembrane 57 - 73 (51 - 80)
 INTEGRAL Likelihood = -6.85 Transmembrane 173 - 189 (169 - 194)
 INTEGRAL Likelihood = -5.79 Transmembrane 94 - 110 (86 - 112)
 INTEGRAL Likelihood = -1.44 Transmembrane 221 - 237 (221 - 238)
 INTEGRAL Likelihood = -1.33 Transmembrane 118 - 134 (118 - 134)
 35 PERIPHERAL Likelihood = 4.72 145
 modified ALOM score: 2.03

*** Reasoning Step: 3

40 ----- Final Results -----
 bacterial membrane --- Certainty=0.4057(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

45 The protein has homology with the following sequences in the databases:

ORF02082(292 - 1053 of 1365)
 EGAD|89511|HP0300(23 - 283 of 285) dipeptide ABC transporter, permease protein (dppC)
 [Helicobacter pylori] OMNI|HP0300 dipeptide ABC transporter, permease protein (dppC)
 50 GP|2313398|gb|AAD07369.1|AE000548 dipeptide ABC transporter, permease protein (dppC)
 [Helicobacter pylori 26695] EIR|D64557|D64557 dipeptide ABC transporter, permease protein -
 Helicobacter pylori (strain 26695)
 %Match = 20.5
 %Identity = 43.4 %Similarity = 63.3
 Matches = 111 Mismatches = 92 Conservative Sub.s = 51
 55

30 60 90 120 150 180 210 240
 P+KCLTCDNDST*LDLGLLINRINYC*RNFFMENNRTFCIDQSKNFRSSSSSTLYANFWNLIFS**FDTIVFYELG*SSV
 MESFR

60 270 300 330 360 402 432 462
 TKVKGELISKRIYFSSLLVLVISAIFAPILSSFPQYVDLSQKILAP-----NNVHLIGTDQLGRDVLRLLYGARY
 ::||| |||||:| || : :|| | :||| |||:|||||
 EPIQQPKNAAVVGAWIVLLVICAIFAPLAPHDPVQNAQRLKPIWEKGNKYLITGDDLRDILSRILYGARI
 65 20 30 40 50 60 70 80

	492	522	552	582	612	642	672	702	
	SFLAIITISIL	ELTGMFGLVGVYQKGL	EMLPIANTIIAR	PSPLSIATVGHIGLGLN	IFALVIVVWVYAKLM				
5	: : : :	: :	: :	: :	: :	: :	: :	: :	
	SLTIGTIVSM	IGVAFPTIL	SLTAYGPGKCTAI	LRIMDIMPAL	PSLILVIVAVL	GPSLITNAML	IGAFVGI	PGPARLV	
	100	110	120	130	140	150	160		
	732	762	792	822	852	882	912	942	
	THVSAKSEK	PEPVINAQIM	GLSVHILK	KHIPPVYQ	PLVLMN	IGNITIL	ISGSPSLG	QVQPNV	TSWGNMLDARG
10	: : :	: :	: :	: :	: :	: :	: :	: :	
	RSVTVGEKEK	YVIAKINGS	SHRLRMCK	VIPNACI	PLIVQT	TINGFAST	VLEAAAL	SFLGLGAQ	PPKPGWAGLMSMQ
	180	190	200	210	220	230	240		
	972	1002	1032	1059	1089	1119	1149		
15	YFRITATMML	SPGIAIFL	TVSFTNLG	DAI-DKIDKWRQ	WNS*K*ENCHYR	*ERSLY*ELIVVK	*IWENR*ELLIVVV		
	: : :	: :	: :	: :	: :	: :	: :		
	YIATAPWLV	PGVITFL	IVNSVL	GDIMDAL	DPKRTS				
	260	270	280						

20 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 24

A DNA sequence (GBSx0021) was identified in *S.agalactiae* <SEQ ID 69> which encodes the amino acid sequence <SEQ ID 70>. This protein is predicted to be peptide ABC transporter, ATP-binding protein.

25 Analysis of this protein sequence reveals the following:

Possible site: 60

>>> Seems to have no N-terminal signal sequence

INTEGRAL Likelihood = -0.32 Transmembrane 161 - 177 (161 - 177)

----- Final Results -----

bacterial membrane --- Certainty=0.1128(Affirmative) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 10027> which encodes amino acid sequence <SEQ ID 10028> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAF73561 GB:AE002315 peptide ABC transporter, ATP-binding

protein [Chlamydia muridarum]

Identities = 86/253 (33%), Positives = 154/253 (59%), Gaps = 2/253 (0%)

Query: 1 METTMEOLEIRKLSLOIGEVPLRDESKIDMGESLTITIGESGSKTLLAKLLVGHIPQG 60

M T+ ++E ++++ ++ S I +SL ++GE+GSGKT ++K ++G +P

Subject: 1 MSKTLK IENLVVAIKESNQR LNVNHL SLTIKQRQSLALVGENGSGKITTVSKAILGFLPDN 60

100

Query: 61 MTVR-GNIFFKGVDLGKLTVKWOKLRGRDIAYLVONPMSMFNPFOKIEAHILETILSHE 119

++ G IF+ G D+ +L+ K++Q +RG+ I+ + QN M P ++ I+ET+ H

Subject: 61 CCIQSGKIFYSGTDITRLSRKEFQSIRGKKISTIFQNAMGTLTPSMRVGTQIIETLRHHF 120

Query: 120 KCSKRVALSKALEWMMKRLNLDDAISLLKKYPFELSGGMLQRINLATILSLDPQVILDEP 179

SK A +KA E + +++++ L+ YPFELSGGM QR+ +A L+ +P++II DEP

Sbjct: 121 VMSKEEAFKARELLVSVHIESPDRCLQLYPFELSGGMCQRVSIATLATNPELTIADEP 180

Query: 180 TSAVDCHNCSTISAILQEL-QNNGKTLITVTHDYQLARDLGGQLLVISEGEVVEQGQTOA 238

++A+D + + + +L+++ QNN L+ +TH+ L +L ++ +I GK+VBQG

Subject: 181 STALDSISQAQVLRVLKQIHQNNNTALLLITHNLALVSELCEMAI IHHGEIVEQGPVHE 240

Query: 239 ILSNPQHNYTKAI 251

+L +P H YT+ L
 Sbjct: 241 LLRSPSHFYTKL 253

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 71> which encodes the amino acid sequence <SEQ ID 72>. Analysis of this protein sequence reveals the following:

Possible site: 55

>>> Seems to have no N-terminal signal sequence

INTEGRAL	Likelihood = -2.50	Transmembrane	168 - 184 (167 - 184)
INTEGRAL	Likelihood = -1.70	Transmembrane	211 - 227 (211 - 227)

----- Final Results -----

bacterial membrane	---	Certainty=0.1999 (Affirmative)	< succ>
bacterial outside	---	Certainty=0.0000 (Not Clear)	< succ>
bacterial cytoplasm	---	Certainty=0.0000 (Not Clear)	< succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 87/232 (37%), Positives = 138/232 (59%), Gaps = 3/232 (1%)

Query: 23 LRDPSCCKIDMGESLTIGESGSKTLLAKLAVGHIPQ-GMTVRGNIFPKGVDLGLK-TVK 80
 +R+ S ++ GE L +GSGSGK++L K G + G G+I ++G +L L T K
 Sbjct: 28 IRNVSLLEVEGEVLAFGVSGSGKGVLTKTFTGMLSENGRIANGSIVTRQGEITDLTKNK 87

Query: 81 QWQKLGRDIAVLVQNPMSMPNPFQKIEAHILETILSHEKCSKRVALSKALEWKKRLND 140
 +W K+RG IA + Q+PM+ +P + I + I E I+ H+K S A AL++M ++ +
 Sbjct: 88 EWAKIRGSKIATIFQDPWTSLSPIKTIGSGQITEVILKHQKVSNAKAKEMALDYANKVGIDP 147

Query: 141 DAISLLKKYPFELSGGMLQRIMLATILSLDPQVILDEPTSAVDCHNCSTISAILQELQN 200
 +A + YPFE SGGM QRL++A L+ P +I DEPT+A+D + I +L+ LQ
 Sbjct: 148 NAKKRFEYDPFYSGGMRQRIVIAIALACRPDLICDEPTIALDVTIQAVIVLLKSLQR 207

Query: 201 NGK-TLITVTHDYQLARDLGGQLLVISEGEVVEQQQTQAILSNPQNTKAL 251
 T+I +THD + + ++ V+ GE+VE G + I +P+H YT +L
 Sbjct: 208 EYHPTIIFITHDLGVVASIADKVAVMVAGEIVEFGTVREIFYDPRHPTWLSL 259

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 25

A DNA sequence (GBSx0022) was identified in *S.agalactiae* <SEQ ID 73> which encodes the amino acid sequence <SEQ ID 74>. This protein is predicted to be peptide ABC transporter, ATP-binding protein. Analysis of this protein sequence reveals the following:

Possible site: 50

>>> Seems to have an uncleavable N-term signal seq

----- Final Results -----

bacterial membrane	---	Certainty=0.0000 (Not Clear)	< succ>
bacterial outside	---	Certainty=0.0000 (Not Clear)	< succ>
bacterial cytoplasm	---	Certainty=0.0000 (Not Clear)	< succ>

A related GBS nucleic acid sequence <SEQ ID 10025> which encodes amino acid sequence <SEQ ID 10026> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:BAB05797 GBS:AP001514 oligopeptide ABC transporter (ATP-binding protein) [Bacillus halodurans]
 Identities = 82/199 (41%), Positives = 130/199 (65%), Gaps = 2/199 (1%)

Query: 19 EQEVLKDCHFHLLKRGELIIGWKSQSGKSSSLARLLIIGLDSPTCGSIYPQG-KIYTFKDGK 77
 +Q++L F + GE +GI+G+SGSGKS+L RL++G++ P G IYF+G K+
 5 Sbjct: 21 KQKILNHLISPECHRGECGLIGESQSGKSLTRLRLLLGIEKPRGHIFYGKMKVEERSVRS 80

Query: 78 AQIILVFQDALSSVNPYFSIEELINEAFYGGKTT-FELCQILSAVGLDGTLYLKYKARQLS 136
 I VFQD SS+NP+P++E + E GKK ++ +L+ VGL +Y K +LS
 Sbjct: 81 GNISAVFQDYTSINPFTVTATMEPLKGGKAASKVDYLKQVGLHPSYKKKYFHELS 140

10 Query: 137 GGQLRQVCIRARALLLPKLIIFDESLSGLDPVTQIKMLRLQKIKRRYELSPINISHDEK 196
 GG+QQRVCIRAR+ +FK I+ DE++S LD Q ++L LL ++KR Y++S++ I+HD +
 Sbjct: 141 GGEVQRVCIRARAISTEPKCIVLDEATSSLDVSIQTQVLDDLILTELKRYIQMSYLFTHIDQ 200

15 Query: 197 ICQICNRVFLIKNGYLVE 215
 IC+R+ + ++G + E
 Sbjct: 201 RAAVICDRIMFRHQIEE 219

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 75> which encodes the amino acid sequence <SEQ ID 76>. Analysis of this protein sequence reveals the following:

20 Possible site: 60
 >>> Seems to have no N-terminal signal sequence

---- Final Results ----
 25 bacterial cytoplasm --- Certainty=0.3195 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

30 Identities = 91/238 (38%), Positives = 137/238 (57%), Gaps = 21/238 (8%)

Query: 1 MKEIFLMIAVCHHVGTQGRQ----EVLKDCHFHLLKRGELIIGWKSQSGKSSSLARLLIIGL 56
 M E + L +H+ TF ++ E +KD H+ +G+I GI+G SG+GKS+L R+V I L
 Sbjct: 1 MNEALIQ--DHDITTFQKRIKRVIRAVKDVTVHINQGDYIGVIGYSGAGKSTLVRINIL 58

35 Query: 57 DSPTCGSI-----YFQGIYTFKDGKQAQ---IILVQ--DALSSVNPYFSIEELINE 103
 +FT G I + CGKI D Q I ++PQ ++ ++ L
 Sbjct: 59 QAPTNGKITVDGVDTFDQGIQLSADALEKQKRRDIGMIFQHFNLMAQKTAKENVAFAIRH 118

40 Query: 104 AFYQK-KITTFELCQILSAVGLDGTLYLKYKARQLSGGQLRVCIRARALLLPKLIIFDES 162
 + K + ++ ++L E VGL Y A QLGGGQ QRV IARAL PKI+I DE+
 Sbjct: 119 SLSKTEKHEKHYELLELVGLSERADNYFA-QLSGGQKQRVARALANDPKILISDEAT 177

45 Query: 163 SGLDPVTQIKMLRLQKIKRRYELSPINISHDEK1CQAIICNRVFLIKNGYLVEDNEFL 220
 S LDP T ++L LLQ++ R+ L+ +M+H+ +I + ICNRV +++NG L+E+ L
 Sbjct: 178 SALDPKTKTQIIALQENRKLGLTIVMITHEMQIVEDIENRVAVMQNGVLEEGSVL 235

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

50 Example 26

A DNA sequence (GBSx0023) was identified in *S.galactiae* <SEQ ID 77> which encodes the amino acid sequence <SEQ ID 78>. This protein is predicted to be UMP kinase (pyrH). Analysis of this protein sequence reveals the following:

55 Possible site: 18
 >>> Seems to have no N-terminal signal sequence

---- Final Results ----
 bacterial cytoplasm --- Certainty=0.1935 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

5 >GP:CB13524 GB:Z99112 uridylyate kinase [Bacillus subtilis]
Identities = 143/238 (60%), Positives = 193/238 (81%)

Query: 2 EPKYQRILIKLSEALAGDKGVGIDIPVQSIKAEIAEVHNSGVQIALVIGGGNLWRGE 61
+PKY+RI++KLSEALAG++G GI+ +QSIK++E+ V++A++V+GGN +
10 Sbjct: 3 KFKYQRILIKLSEALAGDQGGNGINPTVIGSIKQVKEIAELEVEVAVVGGNGVGAET 62

Query: 62 AAEAGMDRVQADYTGMLGTVMNALWADSLQQYGVDTRVQTAIPMOTVAEPYVRGRALRH 121
++ GMGR ADY QML TWMN+L + DGL+ G+ +RVQT+I +H VAREPYR +A+RH
15 Sbjct: 63 GSDLGMDRATADYTKMLATVMNSLALQDSLETIGQSRVQTSIEMRQVAREPYRRKAIKH 122

Query: 122 LEKRNIRIVFGAGIGSPYFSTDTTAAALRAAEIABAILMAKNGVDGVYNADPKKDANAVK 181
LEK R+V+V AG G+PYFSTDTTAAALRAAEI+ IIMAKN VDGYNADP+ED +AVK+
15 Sbjct: 123 LEKRVVIFAAQTGNPYFSTDTTAAALRAAEIADVILMAKNGVDGVYNADPKDASAVKY 182

Query: 182 DELTHVEVIRKGLKIMDATASTISMNDIDLVFNMNETGNIKRVVLGBOIGTIVSNK 239
+ L+++++V+K GL++ND+TAS++ MNDI L+VF++ E GNIR V+GE IGT V K
20 Sbjct: 183 ESLSYLDVLKGLGVNDSTASSLQNDNDIPLIVFSIMEENIKRAVIGBSIGTIVRKG 240

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 79> which encodes the amino acid
25 sequence <SEQ ID 80>. Analysis of this protein sequence reveals the following:

Possible site: 18

>>> Seems to have no N-terminal signal sequence

30 ----- Final Results -----
bacterial cytoplasm --- Certainty=0.1955 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

35 An alignment of the GAS and GBS proteins is shown below:

Identities = 224/242 (92%), Positives = 233/242 (95%)

Query: 1 MEPKYQRILIKLSEALAGDKGVGIDIPVQSIKAEIAEVHNSGVQIALVIGGGNLWRGE 60
+EPKYQRILIKLSEALAG+KGVGIDIPVQ+IAKIAEVH SGVQIALVIGGGNLWRGE
40 Sbjct: 1 VEPKYQRILIKLSEALAGDKGVGIDIPVQAIKAEIAEVHNSGVQIALVIGGGNLWRGE 60

Query: 61 PAAEAGMDRVQADYTGMLGTVMNALWADSLQQYGVDTRVQTAIPMOTVAEPYVRGRALR 120
PAA+AGMDRVQADYTGMLGTVMNALWADSLQ YGVDTRVQTAIPM VAREPY+RGRALR
45 Sbjct: 61 PAADAGMDRVQADYTGMLGTVMNALWADSLQHYGVDTRVQTAIPM+VAREPYIRGRALR 120

Query: 121 HLEKNRIVVFGAGIGSPYFSTDTTAAALRAAEIABAILMAKNGVDGVYNADPKKDANAVK 180
HLEKNRIVVFGAGIGSPYFSTDTTAAALRAAEIABAILMAKNGVDGVYNADPKKDANAVK
45 Sbjct: 121 HLEKNRIVVFGAGIGSPYFSTDTTAAALRAAEIADVILMAKNGVDGVYNADPKKDANAVK 180

Query: 181 FDELTHVEVIRKGLKIMDATASTISMNDIDLVFNMNETGNIKRVVLGBOIGTIVSNK 240
FDELTH EVIRKGLKIMDATAST+SMNDIDLVFNMNE GNI+RVV GE IGTIVSNK
50 Sbjct: 181 FDELTHGEVIRKGLKIMDATASTLSMNDIDLVFNMNETGNIRVVRGHI GTIVSNKV 240

Query: 241 SE 242
+
55 Sbjct: 241 CD 242

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 27

A DNA sequence (GBSx0024) was identified in *S.galactiae* <SEQ ID 81> which encodes the amino acid sequence <SEQ ID 82>. Analysis of this protein sequence reveals the following:

Possible site: 22

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3712(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 28

A DNA sequence (GBSx0025) was identified in *S.galactiae* <SEQ ID 83> which encodes the amino acid sequence <SEQ ID 84>. This protein is predicted to be ribosome recycling factor (frr). Analysis of this protein sequence reveals the following:

Possible site: 34

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3522(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:BA06143 GB:AP001515 ribosome recycling factor [Bacillus halodurans]
Identities = 112/185 (60%), Positives = 149/185 (80%)

Query: 1 MTKIVTKAQERFQSHQSLREFAGIRAGRANASLLDRIQVEYGAFTPLNQLASITVP 60
M+KE++ A++R ++ ++L RE A +RAGRAN ++LDRI VEYGA TPLNQLA+I+VP
Sbjct: 1 MSKEVLNDABQRMKATEALGRBLAKLAGRANPAMLDRIQVEYGAFTPLNQLATISVP 60

Query: 61 EARVLLISPPDKRSIKDIERAINESDLGINPANDGVSIRIVIPALTESTRDLAKEVKKV 120
EAR+L+I PDKRSI DIERAI +SDLG+ P+NDG+VIR+ IP LTEE RRDL K VKK
Sbjct: 61 EARLLVIPCFFDKRSISDIERAIQKSDGLGTPNDGTVIRITIPPLTEERRRDLTKLVKS 120

Query: 121 GENAKIAIRNIRNDAMDHAQKQKKEITDDLSLEKDIQKATDDAVKHIDEMTANKEK 180
E AK+A+RNIRRIA D+ KK++K+ E+TDLL+ + +D+QK TD ++ ID+ KEK
Sbjct: 121 ABEAKVAVRNIRNDANDDLAKRQNDGHLTDLLRRVTSQVQLTKDYITQIDQKAEK 180

Query: 181 ELLEV 185
E++EV
Sbjct: 181 EIMEV 185

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 85> which encodes the amino acid sequence <SEQ ID 86>. Analysis of this protein sequence reveals the following:

Possible site: 21

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

1 bacterial cytoplasm --- Certainty=0.4462 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 5 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 160/185 (86%), Positives = 171/185 (91%)

10 Query: 1 MTKSIVTKAQERFQSHQSLSRFPAGIRAGRANASLLDRIQVEYTGAPTLNQLASITVP 60
 M I+ A+HRF QSHQSLSR+A IRAGRANASLLDRIQV+YTGAPTLNQLASITVP
 Sbjct: 1 MANAIIETAKERFAHQSLSRFYASIRAGRANASLLDRIQVDYTGAPTLNQLASITVP 60

15 Query: 61 EARVLLISPPDKSSINDIERAINESDLGINPANDGSVIRLVIPALTESTRKLAKVEKVV 120
 EARVLLISPPDKSSINDIERA+N SDLGI PANDGSVIRLVIPALTESTRKLAKVEKVV
 Sbjct: 61 EARVLLISPPDKSSINDIERAINASDLGITPANDGSVIRLVIPALTESTRKLAKVEKVV 120

20 Query: 121 GENAKIAIRNIRRDAMDAAKQEKKEITEDDLKSLKDIQKATDDAVKHIDMTANKEK 180
 GENAKIAIRNIRRDAMD+AKQEK KEITED+LK+LEKDIQKATDDA+K ID MTA KEK
 20 Sbjct: 121 GENAKIAIRNIRRDAMDAAKQEKKEITEDDLKSLKDIQKATDDAIKEIDMTANKEK 180

Query: 181 ELLSV 185
 ELL V
 25 Sbjct: 181 ELLSV 185

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 29

A DNA sequence (GBSx0026) was identified in *S. galactiae* <SEQ ID 87> which encodes the amino acid sequence <SEQ ID 88>. Analysis of this protein sequence reveals the following:

Possible site: 44

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

35 bacterial cytoplasm --- Certainty=0.1356 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

40 A related GBS nucleic acid sequence <SEQ ID 10023> which encodes amino acid sequence <SEQ ID 10024> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP: CAB12943 GP:299109 yitL [Bacillus subtilis]

Identities = 107/269 (39%), Positives = 155/269 (56%), Gaps = 6/269 (2%)

45 Query: 42 IATDENKDF-YFIQKDGPTFALSKEBGEHHIGM--VKGPAYTDMQKARLITKETPATR 98
 L D DF YP+ T L SE I+ V+ P Y D Q++ TK +
 Sbjct: 25 LSIHQDTDFGYFLTDGEDTILLHNSMTEDIDRDEVEVPTIVDQGERLATMKIPIISA 84

50 Query: 99 DNYGWIIVTEVRKDLGVFLDTGLPDKQVVSGLDVLPELKLINPKKGDRIYVCLDVDKER 158
 D YGW V+ +D+GVF+D GL K +V+ LP ++NP+EGD+LY L V + R
 Sbjct: 85 DEYGWVEVVDKVEDMGVFDVGL-SKDALVATEHLPPYEDVWPKQKDKLYCMKVTRNGR 143

55 Query: 159 LNALPADPEVQFQFMATPAYNNMNCNNNPFAIVYELKSGTFVYLPENNNLGFTHPSERYSE 218
 ++A PA ++ + T A ++ N+ VYEL SG+PV + ++ FTHPSER E
 Sbjct: 144 MFAKPAFEDIISLPTDASEDLNKKELGTGVYRLIAGSPV-ITDDGRCFTHPSERKEE 202

Query: 219 PRLGQVLDARVIGFREVDRITNLKSPRSFMLENDAQMILTYLESNGQFMTNLNKKSPFE 278
 PRLG + RVI +E D ++NLSL PR ++ DA+ ILTY+ G M +DKS P+

Sbjct: 203 PRLGSRVITGRVIQVKE-DGSVNLSELLPRKQDAMSVDAECILTYMRMRNGAMPYSKSKSQPD 261

Query: 279 EIKATPGISKQGFKKALGGLMKAKIKQD 307
+I+ F +SK FK+ALG LMK K+ Q+

5 Sbjct: 262 DIRERFNMSSKAAFKRALGLHLMKQKVVQE 290

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 89> which encodes the amino acid sequence <SEQ ID 90>. Analysis of this protein sequence reveals the following:

Possible site: 51

10

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

15

bacterial cytoplasm --- Certainty=0.0811 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 235/284 (82%), Positives = 265/284 (92%)

20

Query: 31 MNTLLATVITGLVFDENKDYFYFIQKGGTFALSKSEGEHHIGEMVKGFPAYTDMQQKARIT 90

MN LLATVITGL+ +EN + YFI K+GTF LSK+EGE IG+MV GFAYTD++QKARIT

Sbjct: 1 MNDLLATVITGLIKESNANDYFIHKGGTFPTLSKABGERQIGDMVTGFAYTDIEQKARIT 60

25

Query: 91 TKETPATRDHYGWSVTFVKKDLGVFLDTGLPKQVVSLDLVLEPKELMPKKGDLRYVC 150

TKH +TR YGWS VTEVR+DLGVF+DTG+P+K++VVSLLDLVE+KELMPKKGDL+LY+

Sbjct: 61 TKRIRSTRTSYGWSVTEVRKDLGVFDTGIFNKEIVVSLLDLVEPKELMPKKGDLRYR 120

30

Query: 151 LDVDDKDRLLNALPADPEVQPMATPAYNNMQNMPAIVYRLKLSGTFFVLPENRMGLFI 210

LDVDDKDR+L LPA+PEVQ+MA+PAYNNMQN+MPAIVYRLKL+GTFFVLPENRMGLFI

Sbjct: 121 LDVDDKDRDWGLPAPEVQPMASPAYNNMQNMPAIVYRLKLTGTFFVLPENRMGLFI 180

35

Query: 211 HPSERYSEPRLGQVLDARVIGFREVDRTLNLSLKPRSFEMLENDQMILTYLESNGGFMT 270

H SERY+EPRLGQVLDARVIGFREVDRTLNLSLKPRSFEMLENDQMILTYLESNGGFMT

Sbjct: 181 HSSERYAEPRLGQVLDARVIGFREVDRTLNLSLKPRSFEMLENDQMIVITYLENGGFMT 240

Query: 271 LNDKSSPEEIKATPGISKQGFKKALGGLMKAKIKQDGLTELL 314

LNDKSSPEEIKA+PGISKQGFKKALGGLMKAK+IKQD GTEL+

40

Sbjct: 241 LNDKSSPEEIKASPGISKQGFKKALGGLMKAKIKQDRTGTETLI 284

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 30

A DNA sequence (GBSx0028) was identified in *S.agalactiae* <SEQ ID 91> which encodes the amino acid sequence <SEQ ID 92>. This protein is predicted to be peptide methionine sulfoxide reductase (msrA). Analysis of this protein sequence reveals the following:

Possible site: 33

50

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.0866 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

55

A related GBS nucleic acid sequence <SEQ ID 10021> which encodes amino acid sequence <SEQ ID 10022> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:BA05167 GB:AP001512 peptide methionine sulfoxide reductase
[Bacillus halodurans]
Identities = 102/173 (58%), Positives = 126/173 (71%), Gaps = 2/173 (1%)
Query: 14 ENDMEIRAIFAGGCFWCMVQPFEELOGIRSVLSGYTGSHVENPTYKRVCSKTIGTTHAVEI 73
E+ A FAGGCFWCMV PFEE GI V+SGYTGSH ENPTYKRVCS+ITGH EAV+I
Sbjct: 3 ESKWALATFAGGCFWCMVSPFEEEPGLHQVVSQYTGSHENPTYKRVCSKTIGTTHAVEI 62
Query: 74 IFNPEKISYADLVLYNQTDPDAFGQFEDRGDNRYRPIVYRNKRQRQIAQESKDLQA 133
F+PE Y L+E+YW Q DPTD GQF DRGD+YR IFY +E+Q+Q A SK KL+
Sbjct: 63 SFDPEVFPYEKLLIETWQIDPTDGGQFHDRGDSYRTAIFYHDEQKQAADASKQKLEE 122
Query: 134 SGRFDRPIVTSIEPADTFYPAEDYHQAFYRTNPARYAL--SSARRHAFLENNW 184
SG+F+ PIVT I PA FYPAE+YHQ +++ NP Y + + R AF++++W
Sbjct: 123 SGKFNPIVTRILPAKPFYPAEYHQYKHKKNPFHYKMYRHGSGREAFIKQHW 175

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 93> which encodes the amino acid sequence <SEQ ID 94>. Analysis of this protein sequence reveals the following:

Possible site: 17
>>> Seems to have no N-terminal signal sequence
----- Final Results -----
bacterial cytoplasm --- Certainty=0.0084 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
RGD motif: 89-91

The protein has homology with the following sequences in the databases:

>GP:BA05167 GB:AP001512 peptide methionine sulfoxide reductase
[Bacillus halodurans]
Identities = 98/168 (58%), Positives = 125/168 (74%), Gaps = 4/168 (2%)
Query: 4 AIFAGGCFWCMVQPFEEQAGILSVRSYTGSHLPNPSYEQVCAKTIQHTHAEVLIIFDPKQ 63
A FAGGCFWCMV PFEE+ GI V SGYTGSH NP+Y++VC+ITGH EAV+I FDP+
Sbjct: 9 AIFAGGCFWCMVSPFEEEPGLHQVVSQYTGSHENPTYKRVCSKTIGTTHAVEIIFDPKQ 68
Query: 64 IAYKDLVLYNQTDPDAFGQFEDRGDNRYRPIVYIYTTTERQKEIABQSKANLQASGRFDQ 123
Y+L+E+YWQ DPTD GQF DRGD+YR I+Y E+QK+ A+ SK L+ SG+F+
Sbjct: 69 FPYEKLLEIYWQIDPTDGGQFHDRGDSYRTAIFYHDEQKQAADASKQKLEESGKFN 128
Query: 124 PIVTTIEPAEPFYLAEDYHQGFYKKNP---KRYAQSSAIRHQFLERNW 168
PIVT I PA+PFY AE+YHQ ++KKNP K Y S R F++++W
Sbjct: 129 PIVTRILPAKPFYPAEYHQYKHKKNPFHYKMYRHGSG-REAFIKQHW 175

An alignment of the GAS and GBS proteins is shown below:

Identities = 130/168 (77%), Positives = 148/168 (87%)
Query: 17 MEIRAIFAGGCFWCMVQPFEELOGIRSVLSGYTGSHVENPTYKRVCSKTIGTTHAVEIIFN 76
MEIRAIFAGGCFWCMVQPFEE GI SV SGYTGSH+ NP+Y++VC+KTIGTTHAVEIIF+
Sbjct: 1 MEIRAIFAGGCFWCMVQPFEEQAGILSVRSYTGSHLPNPSYEQVCAKTIQHTHAVEIIFD 60
Query: 77 PEKISYADLVLYNQTDPDAFGQFEDRGDNRYRPIVYIYTTTERQKEIABQSKANLQASGR 136
P++I+Y DLVLYN QTDPTDAFGQFEDRGDNRYRPI+Y E Q++IA++SK IQASGR
Sbjct: 61 PQKIAKDLVLYNQTDPDAFGQFEDRGDNRYRPIVYIYTTTERQKEIABQSKANLQASGR 120
Query: 137 FORPIVTSIEPADTFYPAEDYHQAFYRTNPARYALSSARRHAFLENNW 184
FD+PIVT+IEPA+ FY AEDYHQ FY+ NP RYA SSA RH FLERNW
Sbjct: 121 FDQPIVTTIEPAEPFYLAEDYHQGFYKKNPKRYAQSSAIRHQFLERNW 168

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 31

A DNA sequence (GBSx0029) was identified in *S.agalactiae* <SEQ ID 95> which encodes the amino acid sequence <SEQ ID 96>. Analysis of this protein sequence reveals the following:

Possible site: 55

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2727(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAB13859 GB:Z99114 yoeS [Bacillus subtilis]
Identities = 24/66 (36%), Positives = 42/66 (63%)

Query: 3 KSFYSLWLTQRNPKSNEPVAIDLADYAFDETTFFKHSSDFETVSRYLEDEASFSFNLDTDFD 62

KSFY +L+ R+PK + ++ A+ A+++ +FPK S+D+ +S YLE A + + FD
Sbjct: 2 KSFYHYLLKYRHPKPKDSISEPANQAYEDHSFPKTSIDYHEISSYLEINADYLTMTATFD 61

Query: 63 DIWEDY 68

+ W+ Y

Sbjct: 62 EAWDQY 67

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 97> which encodes the amino acid sequence <SEQ ID 98>. Analysis of this protein sequence reveals the following:

Possible site: 57

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2571(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 59/71 (83%), Positives = 65/71 (91%)

Query: 1 MRKSFYSLWLTQRNPKSNEPVAIDLADYAFDETTFFKHSSDFETVSRYLEDEASFSFNLDT 60

MRKSFYSLWLTQRNPKSNEPVAIDLAD FD+TTFPKH++DFE +SRYLE+ASFSFNL

Sbjct: 3 MRKSFYSLWLTQRNPKSNEPVAIDLADLVFDDTTFFKHINDFELISRYLEDQASFSFNLQ 62

Query: 61 FDDIWRDYLNH 71

FD+IWRDYLN H

Sbjct: 63 FDIWRDYLAH 73

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 32

A DNA sequence (GBSx0030) was identified in *S.agalactiae* <SEQ ID 99> which encodes the amino acid sequence <SEQ ID 100>. This protein is predicted to be antigen, 67 kDa (myosin-crossreactive). Analysis of this protein sequence reveals the following:

Possible site: 14

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -4.57 Transmembrane 28 - 44 (26 - 45)

----- Final Results -----

bacterial membrane --- Certainty=0.2826(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 101> which encodes the amino acid sequence <SEQ ID 102>. Analysis of this protein sequence reveals the following:

Possible site: 26

>>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -4.62 Transmembrane 40 - 56 (38 - 57)

----- Final Results -----

bacterial membrane --- Certainty=0.2848(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

A related sequence was also identified in GAS <SEQ ID 9109> which encodes the amino acid sequence <SEQ ID 9110>. Analysis of this protein sequence reveals the following:

Possible cleavage site: 50

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial membrane --- Certainty= 0.285(Affirmative) < succ>
 bacterial outside --- Certainty= 0.000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty= 0.000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 477/590 (80%), Positives = 542/590 (91%)

```

Query: 3  MRYTNGNFPAFARPKPEVDKKSAYIVGSLAGLAAAVFLIRDQMDQRIRHIFEELEL 62
          M YT+GN+EAPA PRKPEVD+KSAYIVG+GLAGLAAAVFLIRDG M G+RIR+FEELFL
Sbjct: 15  MYTSGNYTAPATPRKPEVDKKSAYIVGTGLAGLAAAVFLIRDGMGMRGIRHIFEELEL 74

Query: 63  SGGSLDGVKRPDGVFVTRGGREMHFECMWDMYRSIPSLVEPDASYLDEFYWLKDDPN 122
          +GSLDGV++P +GFVTRGGREMHFECMWDMYRSIPSL+P ASYLDEFYWLKDDPN
Sbjct: 75  AGSLDGIKPHLGFVTRGGREMHFECMWDMYRSIPSLIETGASYLEDEFYWLKDDPN 134

Query: 123 SSNCRLIHQCNRLSDGDPFLTGTHSKRLVKLWMEETESLGAKTIEVFVSKRFFESNFWT 182
          SSNCRLIHK+GNR++ DG +TLG SKRL+ L+N+TEESLG +TIEE PS++PF+SNFW
Sbjct: 135 SSNCRLIHKGNRVDDGGQYTLKQSKELIHLIMKTESLGDDQTIREFPSSDFKSNFWV 194

Query: 183 YWGFIMAFPEKWSAEMRYAMRFIHHIGGLDPTSLKPKYCNQYDSMVKPII+YLSHSN 242
          YW TMAFPEKWSA+EMRYAMRFIHHI GLDPTSLKPKYCNQYDSMVKPII+YLSHSN
Sbjct: 195 YWATMAFPEKWSAEMRYAMRFIHHIDGLDPTSLKPKYCNQYDSMVKPIATYLSHSD 254

Query: 243 VDVPQFSKVTNISVDPRKNGOKLAKAIHLTVGGEAKTIDLTDFNDPVFVINGSITESTYNGS 302
          VD+QFD+KV+I V+ G+K+AK IH+TV GEAK 1-LTP+D VPFVINGSITE+ YGS
Sbjct: 255 VDIQFDTKV+DIQVQTAGKVKAKTHMTVSGEAKAIHLTEODLVFVINGSITESSYNGS 314

Query: 303 HDTVAKEPDTLGGSMNLWENLAAQSDGFQHPKVYFKDIPKESWFSATATIKDPAIEPFI 362
          H VAKP LGSSMNLWENLAAQSD+QRPKVPFY+D+P ESFVFSATATIK PAIEPFI
Sbjct: 315 HEEVAKPTKALGGSMNLWENLAAQSDGFQHPKVYQDLPASWFSATATIKHAPFPI 374

Query: 363 ERLTHRLDHDGKVTNGGIVIVTDSNNWMSFAIHRQPHFKQKQENETIWIYGLYSNVEGN 422
          ERLTHRLDHDGKVTNGGII+T+TDSNNWMSFAIHRQPHFKQKQENET VMIYGLYSN EGN
Sbjct: 375 ERLTHRLDHDGKVTNGGIIITDSNNWMSFAIHRQPHFKQKQENETIWIYGLYSNVEGN 434

```

Query: 423 YIKKPIKECTGQREITEEWLYHLGVPEMKIHDLSKQYVSTVPMYPIYSYFMPRVKGER 482
 Y+ K IRECTG+ETIEEWLYHLGVPE K I DL+ + Y+TVPMYPIYSYFMPRVKGER
 Sbjct: 435 YVHKIKIECTGQREITEEWLYHLGVPEVDKIDLASQDYINTVPMYPIYSYFMPRVKGER 494

5 Query: 483 PDVIPQSGVNLAFIGNPAESPSRDTVFTEYSIRDTAMEAVYTFPLNIEGVPKVFNSAADI 542
 P VIP GSVNLAFIGNPAESPSRDTVFTEYSIRDTAMEAVY+FLN+ERG+PEVFNSA+DI
 Sbjct: 495 PKVIPDGSVNLAFIGNPAESPSRDTVFTEYSIRDTAMEAVYSFLNVERGIEPEVFNSAYDI 554

10 Query: 543 RVLILGLVYLNDKKSGVEEMDLIPALMRKVMKKIRGTYLELLREAHLL 592
 R IL++ YVINDKK+++DMDLIPAL+ K+G KKI+ T++SELL++A+L+
 Sbjct: 555 RELLKAFYVYLNDKKAIRKMDLIPALIEKIGHKKIKDTFIRELLKDALNM 604

A related GBS gene <SEQ ID 8475> and protein <SEQ ID 8476> were also identified. Analysis of this protein sequence reveals the following:

15 Lipop: Possible site: -1 Crend: 10
 McG: Discrim Score: -19.82
 GVH: Signal Score (-7.5): -1.16
 Possible site: 14
 >>> Seems to have no N-terminal signal sequence

20 ALOM program count: 1 value: -4.57 threshold: 0.0
 INTEGRAL Likelihood = -4.57 Transmembrane 26 - 42 (26 - 45)
 PERIPHERAL Likelihood = 6.79 378
 modified ALOM score: 1.41

25 *** Reasoning Step: 3

----- Final Results -----
 bacterial membrane --- Certainty=0.2826(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 30 bacterial cytoplasm --- Certainty=0.0000(Not Clear)

SEQ ID 8476 (GBS90) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 18 (lane 6; MW 68.5kDa).

The GBS90-His fusion product was purified (Figure 194, lane 11) and used to immunise mice. The
 35 resulting antiserum was used for Western blot (Figure 256A), FACS (Figure 256B), and in the *in vivo* passive protection assay (Table III). These tests confirm that the protein is immunoreactive on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

40 Example 33

A DNA sequence (GBSx0031) was identified in *S.agalactiae* <SEQ ID 103> which encodes the amino acid sequence <SEQ ID 104>. This protein is predicted to be phoH-like protein (phoH). Analysis of this protein sequence reveals the following:

Possible site: 38
 >>> Seems to have no N-terminal signal sequence

45 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.2339(Affirmative) < succ>
 50 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAB14476 GB:Z99117 phosphate starvation-induced protein

[*Bacillus subtilis*]
 Identities = 191/305 (62%), Positives = 241/305 (78%), Gaps = 1/305 (0%)

5 Query: 27 LQHFDMMSLFGSNERHLKLEENLDVTHARTERVQVLGDSEAVETARLTIEALLVLV 86
 L++PD+ +SLFG+ + LKL+E++L++ I RE + V GD +E++ +A + +LL L+
 Sbjct: 12 LKNPDEALSLFGNQDSFLKIMEKDINLINITGETTIYVSGD-DESFOIADRLLGSLAL 70

Query: 67 NRGMTVNTSDVVFTALSMAGNSIDKFVLYEBEIIKDSYGGKPIRVKTGLQKIIYVDSVKNH 146
 +G+ ++ DV+ A+ MA+ ++ F ++YBEEI K++ GK IRVKT+GQ+ YV ++K +
 10 Sbjct: 71 RRGIEISERDVIYAKMAKKELEYFESYBEEITKNAGKSIKRVKTMQREYVAMERN 130

Query: 147 DVVFGIGPAGTGKTLAVTLAVTALKRGQVKRIILTRPAVEAGESLGFLPGDLKEKVDPY 206
 D+VFGIGPAGTGKTLAV AV ALK G +K+IILTRPAVEAGESLGFLPGDLKEKVDPY
 15 Sbjct: 131 DLVFGIGPAGTGKTLAVVKAHVHAKGHIKKIILTRPAVEAGESLGFLPGDLKEKVDPY 190

Query: 207 LRPVYDALYQILGKEQTSRLMREIIEIAPLAYMRGRTLDDAFVILDEAQNNTIMQKMF 266
 LRP+YDAL+ +LG + T RLNER IIEIAPLAYMRGRTLDDA+VILDEAQNNT QMKMF
 20 Sbjct: 191 LRPLDALHDLVGAHTERLMERGIIIEIAPLAYMRGRTLDDAYVILDEAQNNTTFAQMKMF 250

Query: 267 LTRLGFNSKMIIVNGDVSQIDLPRNVKSGLIDAVEKLENIKKIDFIHLSAKDVVRHPVVAE 326
 LTRLGF+SKMI+ GDVSQIDLPK VKSLG A E L+ I I I L DVVRHP+VA+
 25 Sbjct: 251 LTRLGFSKMIITGDVSQIDLPRGVKSLAVAKEMLEKIDGISMIELDQDVVRHPVNAK 310

Query: 327 IINAY 331
 II AY
 30 Sbjct: 311 IIEAY 315

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 105> which encodes the amino acid sequence <SEQ ID 106>. Analysis of this protein sequence reveals the following:

30 Possible site: 42

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.85 Transmembrane 54 - 70 (54 - 70)

35 ----- Final Results -----
 bacterial membrane --- Certainty=0.1341(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

40 An alignment of the GAS and GBS proteins is shown below:
 Identities = 274/322 (85%), Positives = 298/322 (92%)

Query: 18 LQESYIEITLQHPDDMMSLFGSNERHLKLEENLDVTHARTERVQVLGDSEAVETARL 77
 LQESYI+ITL HPDD++LFGSNERHLKLE +L VI+HARTERVQV+GD BEAVE ARL
 45 Sbjct: 1 LQESYIDITLTHPDVLAFLFGSNERHLKLEINLGVTHARTERVQVLGDSEAVETARL 60

Query: 78 TIEALLVLVNRGMTVNTSDVVFTALSMAGNSIDKFVLYEBEIIKDSYGGKPIRVKTGLQK 137
 TI+ALLVLV RGM VNTSDVVFTALSM++ ID+F+ALYEBEIIKDSYGGKPIRVKTGLQK
 50 Sbjct: 61 TIKALLVLVGRGMVNTSDVVFTALSMAGNSIDQF+ALYEBEIIKDSYGGKPIRVKTGLQK 120

Query: 138 IYVDSVKRHDVVFGIGPAGTGKTLAVTLAVTALKRGQVKRIILTRPAVEAGESLGFLPG 197
 YVDSVK HDVVFG+GPAGTGKTLAVTLAVTALKRGQVKRIILTRPAVEAGESLGFLPG
 55 Sbjct: 121 TYVDSVKRHDVVFGIGPAGTGKTLAVTLAVTALKRGQVKRIILTRPAVEAGESLGFLPG 180

Query: 198 DLKEKVDPYLRPVYDALYQILGKEQTSRLMREIIEIAPLAYMRGRTLDDAFVILDEAQN 257
 DLKEKVDPYLRPVYDALY ILGKEQT+RLMER++IIEIAPLAYMRGRTLDDAFVILDEAQN
 60 Sbjct: 181 DLKEKVDPYLRPVYDALYHILGKEQTSRLMERDVIRIAPLAYMRGRTLDDAFVILDEAQN 240

Query: 258 TTIQMKNFLTRLGFNSKMIIVNGDVSQIDLPRNVKSGLIDAVEKLENIKKIDFIHLSAKD 317
 TTIQMKNFLTRLGFNSKMIIVNGD SQIDL+NVKSGLID+KIL+ IK+IDF++ SAKD
 65 Sbjct: 241 TTIQMKNFLTRLGFNSKMIIVNGDTSQIDLPRNVKSGLIDATQLGLQIKIDFVYFGAKD 300

Query: 318 VVRHPVVAEIIINAYSDSESHK 339
 VVRHPVVA+II AY S K
 65 Sbjct: 301 VVRHPVVAIIKAYETSSEBMK 322

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 34

- 5 A DNA sequence (GBSx0032) was identified in *S.agalactiae* <SEQ ID 107> which encodes the amino acid sequence <SEQ ID 108>. Analysis of this protein sequence reveals the following:

Possible site: 30

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.0275 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

20 Example 35

A DNA sequence (GBSx0033) was identified in *S.agalactiae* <SEQ ID 109> which encodes the amino acid sequence <SEQ ID 110>. This protein is predicted to be MutT/nudix family protein. Analysis of this protein sequence reveals the following:

Possible site: 46

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2383 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>SP:RAF09597 GB:AB001864 MutT/nudix family protein [Deinococcus radiodurans]
Identities = 49/136 (36%), Positives = 69/136 (50%), Gaps = 8/136 (5%)

Query: 5 YISYIRSKVGHETIFITYSGHILTDGKGRVLLQLRADKNSWGLGGCMELGESSVDTLKR 64
Y+S +R+ GH + +L D GRVLLQ R D WGI+GG +E GR + R
Sbjct: 6 YLSELRVAVWGHRLPAAGSVLLQDETGRVLLQRRGDDGQWGLGGCLRPGRDFLIAAR 65

Query: 65 EFPEETGLRVEPIKLLNVY-----TWPDQSYFNGDKACQVGFYIRVSCPKPVNIEGFEN 116
E EETGLR +R L + F YENGD+ VG E + P + +
Sbjct: 66 ELLEETGLRCPNLRLPLSLSGLVSGPQFWHRYFNGDEVYLVGLRTETGVPAARLTDACPD 125

Query: 119 E--ETLQLDYFSKQDV 132
+ ETL+L +F+ +D+
Sbjct: 126 DGSETLELRWFALEDL 141

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 111> which encodes the amino acid sequence <SEQ ID 112>. Analysis of this protein sequence reveals the following:

Possible site: 61

-88-

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

5 bacterial cytoplasm --- Certainty=0.4375 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 93/157 (59%), Positives = 123/157 (78%)

10 Query: 1 MQQDYISYIRSKVGHETIFATYSGGILTDGKGRVILQLRADKNSWGIIGGOMELGESSVD 60
 M QDYISYIRSKVGH+ I L ++GGILT+ G+VL+QLR DK +W I GG MELGESS++
 Sbjct: 16 MPQDYISYIRSKVGHDKIILNFAGGILTRDDGKVLMLQLRGDKKKTWTIPGGIMELGESSLE 75

15 Query: 61 TLKREFFESTGLRVEPIRLNLVYTNFQDSYPWQDKAQTGVPFIYEVSCPKPVNIEGFHNEE 120
 T KREF ESTG+ VE +RLNLVYT+ F++ YPNKD QT+ FIYE++ + I+ FHNEE
 Sbjct: 76 TKKREFLESETGIEVEAVRLNLVYTHFEVYYPNGDAVQTIIVFIYELTAVSDMAIDFNHNEE 135

20 Query: 121 TLQLDYFSKEDVQMITIVNEQHQLILEYFSQTFQMG 157
 TL+L +FS E++ + V+ +H+L+L+EYFS +F MG
 Sbjct: 136 TLKLQFFSHREETABLESVAKHRLNLEEFSDSFAMG 172

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

25 Example 36

A DNA sequence (GBSx0034) was identified in *S.agalactiae* <SEQ ID 113> which encodes the amino acid sequence <SEQ ID 114>. Analysis of this protein sequence reveals the following:

Possible site: 13

30 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

35 bacterial cytoplasm --- Certainty=0.3690 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

40 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 37

A DNA sequence (GBSx0035) was identified in *S.agalactiae* <SEQ ID 115> which encodes the amino acid sequence <SEQ ID 116>. Analysis of this protein sequence reveals the following:

Possible site: 25

45 >>> Seems to have a cleavable N-term signal seq.

----- Final Results -----

50 bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:ANG05249 GB:AE4612 hypothetical protein [Pseudomonas aeruginosa]
Identities = 70/254 (27%), Positives = 127/254 (49%), Gaps = 2/254 (0%)

Query: 2 KITLHGAVETLLITLYIAKDKAMAKHPILINDKSLAIYHQIEYDFKDFKNSRASFYATLA 61
+ITL G +TLLITLY +A D+ II+D+ + V QI+DF+ + + A
Sbjct: 5 RITLTGKQETLLITLYIAKDLSRLLDSILITDFAREAVRQIDYDFPSRLVKGKNEALAM 64

Query: 62 RIRVDRKKKKKIFKFFMNSQILSIGCGLDTRFRVD+NGQIRVRLDLFVMEIRKLFFE 120
R D+ ++P+ +F QI+I+GCGLD+R VDD ++ W+LD FVW+I+R+ +
Sbjct: 65 RSHYFDQCRFPL+RHFFGQVILNIGCGDLSRIVRVDPVALEFDFLDFYFVMDIRERLY 124

Query: 121 EERVINTIAKSLAEDTWTREVNQNPAPFLVISGVMFLKSDVETFLHLILINSFQFMA 180
++D+ + D+ P+I++BQ+ ++I+Z V + L+
Sbjct: 125 PRAGAYALRHSGVDGGLGVPPREPAVLNAGKAMPYLRSSQVRRLRVLVDHLSGEL 184

Query: 181 QFDLCHKEMKNGKHQDITVKYMTDFQPGITDGHISVLDLDEPKLQINLINFIDEMSKFEL 240
FD + I + + + + + I D E+ P L I + I D +L
Sbjct: 185 LFDGVRGRIIGLHLTPPELTAQVNNISIDDPELRERHFAIRFIEVTVYDQDVAK 244

Query: 241 -GTLRSLLPITIRKF 253
+ R +LP F
Sbjct: 245 POSSRLMLPIYNGF 258

```

No corresponding DNA sequence was identified in *S.pyogenes*.

25 A related GBS gene <SEQ ID 8477> and protein <SEQ ID 8478> were also identified. Analysis of this protein sequence reveals the following:

```
Lipop: Possible site: -1  Crend: 9
McG: Discrim Score:      0.37
GVH: Signal Score (-7.5): -0.97
Possible site: 25
>>> Seems to have a cleavable N-term signal seq.
ALOM program count: 0 value: 4.35 threshold: 0.0
PERIPHERAL Likelihood = 4.35 143
modified ALOM score: -1.37
```

*** Reasoning Step: 3

----- Final Results -----

```
bacterial outside --- Certainty=0.3000(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>
```

The protein has homology with the following sequences in the databases:

```

27.6/51.6% over 253aa
      GP|9947849| hypothetical protein Insert characterized
Pseudomonas aeruginosa

ORF02095 (304 - 1059 of 1404)
GP|9947849|gb|AAG05249.1|AE004612_3|AE004612|5 - 258 of 275) hypothetical protein
[Pseudomonas aeruginosa]
%Match = 11.6
%Identity = 27.6 %Similarity = 51.6
Matches = 70 Mismatches = 121 Conservative Sub.s = 61

255      285      315      345      375      405      435      465
E*YTRNPVLEIQISKNSIKESRKKITILGVASTLITLITLIRAKDAMAKIPILAAQKSLAIVQIEYDFKFNSEAS
      |||  |||  |||||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||
      MFGHRIITLQSGKQITLITLAKALDSKLDSILHDFAPFAARQIIDFDFPSVALGKGN
              10          20          30          40          50

495      525      555      585      612      642      672      702
PYATLARXRVMDRIKKFPIKFNENFNSQILSGCGLDTPFRVFN-QGIRVWNLDPWEIRKILFEEHIERINLAKSALD
      |||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||  |||
      ERLALMRSHYFDQACKELGRHPFQVNLGCLDSIRYKLVDPPEALPVPVLDLTPYEWDLRERYLPFRAGYALRHSVD

```

```

      70      80      90      100      110      120      130
5      732      762      792      822      852      882      912      942
      EIVTREVNPQNAFLIVSEGVLMFLAKEDDVETFLHILINSFSQPMADFLCHKEMINKGKHDTVKYMDTFQFGITDGH
      : : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : |
      DDGKWLQGVPRPRPALVLAEGMLPYLRESQVRKILVERLVDHLGSGELLFDGYGRGLIMLLRLVPLPLETGAQVHNSIDDR
      150      160      170      180      190      200      210

10     972      1002      1029      1059      1089      1119      1149      1179
      EIVDLDPKIKQINLINFDEMSKPELG-TLRSLLPTIRKFNCLGVVETKASEKK*QKSIYIKRHSCKCFVILVIAFVAL
      | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : | : : : |
      ELERHHPALRFIREVIDYDPQWAKLPQSSRIMLPYNGFAFLRMERLIRYMERV
      230      240      250      260      270

```

- 15 SEQ ID 8478 (GBS176) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 36 (lane 5 & 6; MW 30kDa). It was also expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 41 (lane 7; MW 55.4kDa).

The GBS176-GST fusion product was purified (Figure 117A; see also Figure 202, lane 5) and used to immunise mice (lane 1+2 product; 13.5µg/mouse). The resulting antiserum was used for Western blot (Figure 117B), FACS (Figure 117C), and in the *in vivo* passive protection assay (Table III). These tests confirm that the protein is immunoreactive on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

25 Example 38

A DNA sequence (GBSx0036) was identified in *S.galactiae* <SEQ ID 117> which encodes the amino acid sequence <SEQ ID 118>. Analysis of this protein sequence reveals the following:

```

Possible site: 32
30 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.3712 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
35      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 10019> which encodes amino acid sequence <SEQ ID 10020> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```

40 >GP:AC38046 GB:AF000954 No definition line found [Streptococcus mutans]
      Identities = 140/164 (85%), Positives = 157/164 (95%)

Query: 1 MYVEMIDETQVSKDIKKQTLDDLEFAAQKTGKEMAVTFVYNERSSHELALKYRDTOR 60
      MYEMIDET QVSR IK QTLD+LEFAAQKTGK+KEMAVTFVYNERSSHELAL+YRDT+R
45 Sbjct: 1 MYEMIDETQVSEGIKNQTLDDLEFAAQKTGKEDKEMAVTFVYNERSSHELALKYRDTOR 60

Query: 61 PTDVISLEYKPEVDISFDEEDLARNPRIARMLADPDSTIGELFISIDKAKQAKKYGH5Y 120
      PTDVISLEYKDE +SFDEEDLA+P+LAE+L+FD+YIGELFIS+DKA+QA+KYGH5+
50 Sbjct: 61 PTDVISLEYKPESSLSFDEEDLADDEDLAKVLTETFDAYIGELFISVDKAREQAQYGH5F 120

Query: 121 EREMGLAVRGFLHTNGYDEYTPSEKEMFSLQSEITLAYGLKR 164
      EREMGLAVRGFLHTNGYDHTTP+SEKEMFSLQSEITL+AYGLKR
Sbjct: 121 EREMGLAVRGFLHTNGYDHTTPQSEKEMFSLQSEITLAYGLKR 164

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 119> which encodes the amino acid sequence <SEQ ID 120>. Analysis of this protein sequence reveals the following:

Possible site: 49

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1145(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 138/165 (83%), Positives = 153/165 (92%)

Query: 1 NYVEMIDETGQVSEIDIKQTLDLLEFAAQTGKGNKEMAVTFVTKRSHEINLEYRDTDR 60
NY+EMIDETGQVS++I +QTLDLL FAAQTGKS KEM+VTFVTKRSHEINLEYRDTDR
Sbjct: 18 NYIEMIDETGQVSEIIMEQTLDLLEFAAQTGKKEKEMSVTFVTKRSHEINLEYRDTDR 77

Query: 61 PTDVISLEYKPEVDISFDEEDLAENPELARMLEDFOSYIGELFISIDKAKRQAREYGHYS 120
PTDVISLEYKPE I F +EDLA +P LARM+ +PD+YIGELFISIDKA+EQ++EYGH+
Sbjct: 78 PTDVISLEYKPE+TFLFSQSDLAADPSLARMMAEFDAYIGELFISIDKARQSQEYGHFS 137

Query: 121 EREMGFLAVHGFLHNGYDHYT+PEERKQMFSLQEELLTAYGLKRRQ 165
EREMGFLAVHGFLHNGYDHYT+ BEEKEM+LQEEILLTAYGL RQ
Sbjct: 138 EREMGFLAVHGFLHNGYDHYT+PEERKQMFSLQEELLTAYGLTRQ 182

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 39

A DNA sequence (GBSx0038) was identified in *S.agalactiae* <SEQ ID 121> which encodes the amino acid sequence <SEQ ID 122>. This protein is predicted to be phosphoglycerate dehydrogenase (serA) (serA). Analysis of this protein sequence reveals the following:

Possible site: 59

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2817(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AB99020 GB:U67544 phosphoglycerate dehydrogenase (serA)
[Methanococcus jannaschii]

Identities = 82/232 (35%), Positives = 132/232 (56%), Gaps = 14/232 (6%)

Query: 3 ENPDYAIRSQNLIHQDQF---PSNLKAIARAGAGTNNIPIEASAGQIVVFNTPGANNA 59
++D ++RS ++D LK I RAG G +NI +E A+ +GI+V N P A++ +
Sbjct: 40 KDADVLVVRSQTKVTRDVIEKAKKLVYGRAGVGVNDIVBAATERGILTVVRAPDASSIS 99

Query: 60 VKEAVIALLLSARDYLGANRWVNTLTGTDIPKQIEAGKAFAGNIEAGKGLVIGLIGAI 119
V E + +L +AR N T K+ E +K F G +E +GK LGVIGLG I
Sbjct: 100 VAEIYMGIELMAAAR-----NIPQNTASLKRGEWDRKRFGKSIELYGKTLGVIGLGRI 150

Query: 120 GARIANDARRLGMTVLGYDPPYVSIETAWNISSHVQRKEIKDIFETCDYITIHVELTINET 179
G + + A+ GM ++GYDPY+ E A + + V+ V +I ++ +D+IT+HVPLT +T
Sbjct: 151 GQQVVKRAKAPGMNIIIGDYPYIKVAVSSMG--VELVDDINELCKRADFTYLLHVPLTPKT 208

Query: 180 KHTFDKAFSINKGGTTIINFARAEIWNQELFEAIETGVVKKYITDPGDKR 231
 +H + ++MKK I+N AR L++ + L+EA++ G ++ D ++E
 Sbjct: 209 RHITGREQIALNKGNIAIVNCARGGLIDEKALYKALKEGKIRAAALDVPEE 260

- 5 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 123> which encodes the amino acid sequence <SEQ ID 124>. Analysis of this protein sequence reveals the following:

Possible site: 52

>>> Seems to have no N-terminal signal sequence

10

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2384 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

15

An alignment of the GAS and GBS proteins is shown below:

Identities = 52/198 (26%), Positives = 93/198 (46%), Gaps = 14/198 (7%)

20

Query: 24 LKAIARAGAGTNNIPIEBASAQGIVVFMTPGANANAVKEAVIAALLLSARDYLGANRWVN 83
 +K IA+ A + ++A+ I++ N P ++ E + +L R
 Sbjct: 70 IKQIAQHSASVDYNNLDELATENDIIITNVPSPESLAEFTVTIVLNLRIHV----- 121

25

Query: 84 TLTGIDIPKQIEAGKKAFAGNEIAGKKLGVILGAGIARIANDARILGMITVLGYDPYVSI 143
 L ++ KQ G + ++IG G IG A + G V+GYD Y S
 Sbjct: 122 ELIRENVKKQNTWGLPIRGVILGDMTVAIIGTGRIGLATAKIPKSGFCKVVGDIYQS- 180

30

Query: 144 ETANNISSHVQRVKE-IKDIFETCDYITIHVPLTNETKHTFDKAFSINKGGTTIINFAR 202
 +A + + + V+E IKD D +++E+P T E H F++ F KKG ++N AR
 Sbjct: 181 DAAKAVLDYKESVEEAIKD----ADLVSLHMPPTAENTHLSNLFKSFKKGAILMNNAR 236

Query: 203 AELVNNQELFEAIETGVV 220
 ++ Q+L +A++ G++
 Sbjct: 237 GAVIETQDLILDALDAGLL 254

- 35 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 40

- A DNA sequence (GBSx0039) was identified in *S.galactiae* <SEQ ID 125> which encodes the amino acid sequence <SEQ ID 126>. This protein is predicted to be alpha-glycerophosphate oxidase. Analysis of this protein sequence reveals the following:

Possible site: 50

>>> Seems to have no N-terminal signal sequence

45

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2067 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

- 50 The protein has homology with the following sequences in the GENPEPT database:

>GP:ABC34740 GB:U94770 alpha-glycerophosphate oxidase [Streptococcus pneumoniae]
 Identities = 24/49 (48%), Positives = 37/49 (74%)

55

Query: 1 MLFMRKNDLSIQVIDEMAKHYQNSDQDKTFYRELHETLKDNLDLAL 49
 MLFMRD+LDS++PV+DEM + Y W++K K Y ++ L +NDLA L
 Sbjct: 558 MLFMRDLSLDSIVEVLDERGFRPDWTEEEKATYTRADVEAALANNLDLAE 606

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 127> which encodes the amino acid sequence <SEQ ID 128>. Analysis of this protein sequence reveals the following:

```

Possible site: 40
>>> Seems to have no N-terminal signal sequence
5  INTEGRAL      Likelihood = -1.81      Transmembrane      20 - 36 ( 20 - 36 )

----- Final Results -----
      bacterial membrane --- Certainty=0.1723 (Affirmative) < succ>
10     bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the databases:

```

>GP:AA034740 GB:U94770 alpha-glycero-phosphate oxidase [Streptococcus pneumoniae]
Identities = 462/607 (76%), Positives = 539/607 (88%)
15  Query: 1  MEFSRETRRLALQKQCRDLLOLLIIGGGITGAGVALQAAASGLTGLIEMQDFAQTSSR 60
      MEFS++TR L+++KQCR LOLLIIIGGITGAGVALQAAASGL+TGLIEMQDFA+GTSSR
      Sbjct: 1  MEFSKTRTELSIKKQCRDLLOLLIIGGGITGAGVALQAAASGLTGLIEMQDFAQTSSR 60

20  Query: 61  STKLHVHGLRLYLKQFDVEVVSVDTSERAVVQQLAPIHPKPDFMLLPVYDEFGSTFMSFEL 120
      STKLHVHGLRLYLKQFDVEVVSVDTSERAVVQQLAPIHPKPDFMLLPVYDE G+TFS+FEL
      Sbjct: 61  STKLHVHGLRLYLKQFDVEVVSVDTSERAVVQQLAPIHPKPDFMLLPVYDEGATGTSFEL 120

25  Query: 121 KVAMLDYLLAGVSNTPANKVLTKEEVLKREPDLKQEGLLGGVYLDFRNNDARLVIN 180
      KVAMLDYLLAGVSNTP ANKVL+K++VL+R+P+LK+EG+GGVYLDFRNNDARLVIN
      Sbjct: 121 KVAMLDYLLAGVSNTPANKVLSKQVLERQPNLKEGLVGGVYLDFRNNDARLVIN 180

30  Query: 181 IKRANRDGALIAHVKAEDFLDDNGKILGVKARDLLSDQEIIRAKLVINTGFWSDIEI 240
      IKRAN+DGALIA+HVKA E FL D++GKI GV ARDLL+DQ IKA+LVINTGFWSD++
      Sbjct: 181 IKRANQDGAIIANHVKAEGFLPDSGKIVGVVARDLLTQVFIRAKLVINTGFWSDIKV 240

35  Query: 241 RQFSHGQPIHQMRPTKGVHLVVDKQLVPVQPVYDTGLDGRNVFVLPREKTYFGTT 300
      R S+KG QMRPTKGVHLVD K+ VQPVY DTGL DGRNVFVLPRE KTYFGTT
      Sbjct: 241 RNLNKGTFQFSQMRPTKGVHLVVDSSKIKVQPVYDTGLDGRNVFVLPREKTYFGTT 300

40  Query: 301 DTDYTGDLHPKVTQEDVDYLLGVVNRFPNANVTIDDISSWAGLRPLINGNSASDYG 360
      DTDYTGDLHP+VTQEDVDYLLG+VNNRFP +N+TIDDISSWAGLRPL+GNSASDYG
      Sbjct: 301 DTDYTGDLHPKVTQEDVDYLLGVVNRFPESNTITIDDISSWAGLRPLINGNSASDYG 360

45  Query: 361 GNSGKVSDDSDHLVDITVKAYINHEDSRZAVEKAIQVETSTSEKLDPSAVRSGSSFER 420
      GN+G +SD+SFD+L+ TV++Y++ E +R E VA ++ ++E+STSEK LDPSAVRSGSS +R
      Sbjct: 361 GNGGTISDESFDNLATVSEYLSKEKTRDVSASVK.ESSSTSEKHLDPASVRSGSLK 420

50  Query: 421 DENGFLTLAGEKITDYRMAAGALTOIITLKEEFGKSFKLINSKTYPVSGEINPNVND 480
      D+NGL TLAGKITDYRMAAGBA+ ++ ILK EF +SFKLINSKTYPVSGE+NPANVD
      Sbjct: 421 DNGLLTLAGEKITDYRMAAGAMRVVDILKAFDRSFKLINSKTYPVSGELNPANVD 480

55  Query: 481 SEIEAYAQGLTSLSLMDDARYLANLYGSNAKVPFALTRQLTAAGLSLAETLSLHYAMD 540
      SEIEA+AQGL GL +A YLANLYGSNAKVPFAL L A GLSLA+TSLSLHYAM
      Sbjct: 481 SEIEAPAQGLVSRGLDSKRAHYLANLYGSNAKVPFALASLEQAQGLSLADTSLSLHYAM 540

60  Query: 541 YEMALKPTDYFLRKTNHLLFMRDSLDLIDPVINEMAKHPFWSQERVAQEDDLRRVIAD 600
      E+AL P D+ LRKTNH+LFMRDSLD++++PV++EM + ++N+++E+ D+ +A+
      Sbjct: 541 NELALSPVDVFLLRKTNHLLFMRDSLDSEIVPELDGMRFDWTESEKATYRADVEALAN 600

65  Query: 601 NDLSALK 607
      NDL+ LK
      Sbjct: 601 NDLEALK 607

```

60 An alignment of the GAS and GBS proteins is shown below:

```

Identities = 29/49 (59%), Positives = 41/49 (83%)

Query: 1  MLFMRNLDLSIQPVIDEAKHYQWSDQKTFYEEELHETIKDNDLAL 49
      +LFMRD+LD+LI PVI+EMAKH++WSDQ++ E++L + DNDL+AL

```

Sbjct: 558 LLFMRDSLALALDPVINEAKHPFWSQRRVAQEDDLRRVADNDLSAL 606

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

5 Example 41

A DNA sequence (GBSx0040) was identified in *S.galactiae* <SEQ ID 129> which encodes the amino acid sequence <SEQ ID 130>. Analysis of this protein sequence reveals the following:

Possible site: 40

10 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

15 bacterial cytoplasm --- Certainty=0.1011 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:BA06309 GB:AF001516 unknown conserved protein [Bacillus halodurans]
Identities = 70/160 (43%), Positives = 106/160 (65%), Gaps = 3/160 (1%)

20 Query: 5 TRPTTDKVGGAIFNMIGPFFEGGRVLDLPSGSGSLAIEAISRGMDQAVLVEKORRAQVVI 64
TRPTTDKVK AI FNMIGPFF+GG LDI+ GSG L IEA+SRG+++ V++ +RA I
Sbjct: 21 TRPTTDKVGGAIFNMIGPFFDGGIGLDLPSGSGSLGIEALSRGVEMIFVDQCKRAIETI 80

25 Query: 65 QENIAMTKSPBQQLLMKANRALEQLTQ---FDLVLLDPFYAKEIVKIQIMDSKGL 121
++N++ ++ +A RAL+ LT+ F V LDPPYAK+ I + I+ + GL
Sbjct: 81 KQNLSHCGLEGRASVYRNDAKRALQVLTRGIVFAYVFLDPFYAKQITRNDLAILANHGL 140

30 Query: 122 LGDDIMIACETDKSVLPEEIASFGIWKQKIYGISKVTIVY 161
L + ++ CE D+ LP++I K++ YG + +T+Y
Sbjct: 141 LEBGGVVVCEHRTIMLPDQIEAYVHKKEITYGDTMITTY 180

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 131> which encodes the amino acid sequence <SEQ ID 132>. Analysis of this protein sequence reveals the following:

35 Possible site: 58

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

40 bacterial cytoplasm --- Certainty=0.3814 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

45 Identities = 111/160 (69%), Positives = 136/160 (84%)

Query: 3 RTRPTTDKVGGAIFNMIGPFFEGGRVLDLPSGSGSLAIEAISRGMDQAVLVEKORRAQV 62
+ TRPT+DKV+GAIFNMIGP+? GORVLDLP+GSG LAIEA+SRGM AVLVEK+R+AQ
Sbjct: 19 KITRPTSDKVRGAIFNMIGPFFNGGRVLDLPSGSGSLAIEAVSRGMSAAVLVEKORRAQA 78

50 Query: 63 VIQENIAMTKSPBQQLLMKANRALEQLTQFDLVLLDPFYAKEIVKIQIMDSKGL 122
+IQ+NI MTK+ +F LLKMEA RA++ LTG+FDLV LDPPYAKE IV + + K LL
Sbjct: 79 IIQCNIIIMTKAENRPTLLMKRAERAI DCLTGRFDLVLDPPYAKETIVATIEALAAKGL 138

55 Query: 123 GDDIMIACETDKSVLPEEIASFGIWKQKIYGISKVTIVY 162
+ N+ CETDK+V LP+EIA+ GIWK+KIYGISKVTIVY
Sbjct: 139 SBQVMVVCETDKTIVLLPKETATLGWKEKIYGISKVTIVY 178

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 42

A DNA sequence (GBSx0041) was identified in *S. galactiae* <SEQ ID 133> which encodes the amino acid sequence <SEQ ID 134>. This protein is predicted to be lipopolysaccharide core biosynthesis protein kdtB (kdtB). Analysis of this protein sequence reveals the following:

```
Possible site: 17

>>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.1937 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

The protein has homology with the following sequences in the GENPEPT database:

```
>GP:BAB13272 GB:AP001119 lipopolysaccharide core biosynthesis
      protein kdtB [Buchnera sp. APS]
      Identities = 56/149 (37%), Positives = 94/149 (62%)

Query: 1  MTKKALFTGSDPVTNGHLDIIERASYLFDHVIYIGLFYNLEKQGYFSICRKNMLEAIR 60
      M K A++ G+FD+T GHLDII RA+ +FD + I + N K+ P+++ R ++ +
Sbjct: 1  MNKTAIYPGTFDPIITGHLDIITRAIKIFDSITIAISNNFKKPIFNKERIELTRKVTIL 60

Query: 61  QFQNVSVLVAQDRILAVDLAREVGAKYFVGRGLNSQDFDYERANLEFFNQQLADDIETVYLS 120
      KNV ++ + L +LA++ A +RG+R DFDYE L NKG+ D+++++L
Sbjct: 61  HLQNVKKILGFNDLLANLAKKERANILIRGVRTITFDYDIKLAALNQLYDLDISIFLL 120

Query: 121 TSPSLSPFISSSRIRLEIHFKAASKVFPVK 149
      +S +S ISSS ++E+ +K +KP++PK
Sbjct: 121 SSKEVSFPISSSPVKEIAKYKGIKPYLPK 149
```

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 135> which encodes the amino acid sequence <SEQ ID 136>. Analysis of this protein sequence reveals the following:

```
Possible site: 61

>>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.1862 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

An alignment of the GAS and GBS proteins is shown below:

```
Identities = 88/161 (54%), Positives = 124/161 (76%)

Query: 1  MTKKALFTGSDPVTNGHLDIIERASYLFDHVIYIGLFYNLEKQGYFSICRKNMLEAIR 60
      +TK L+TGSDPVTNGHLDI++RAS LFD +Y+G+F N K+ YF +E RK ML +A+
Sbjct: 2  LTKIGLYTGSDPVTNGHLDIVKRASGLDQIYVGIFDNPTKSYFKLEVRQMLTQALA 61

Query: 61  QFQNVSVLVAQDRILAVDLAREVGAKYFVGRGLNSQDFDYERANLEFFNQQLADDIETVYLS 120
      F NV V+ + +RLA+D+A+3+ + +RGLRN+ DF+YE NLE+FN LA +IETVYL
Sbjct: 62  DPTNVIVVTISHERLAIDVACELRVTHLIRGLRNATDFEYERANLEYFNHLLAANIETVYLI 121

Query: 121 TSPSLSPFISSSRIRLEIHFKAASKVFPVKSVVREVERKSEE 161
      + +SSSR+RELIIHF++S++ VP+SV+ +VERN+E+
Sbjct: 122 SRNKVQALSSSRVRELIHFQSSLEGLVQSVIAQVEKNEK 162
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 43

A DNA sequence (GBSx0042) was identified in *S.agalactiae* <SEQ ID 137> which encodes the amino acid sequence <SEQ ID 138>. Analysis of this protein sequence reveals the following:

Possible site: 15

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1126 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 44

A DNA sequence (GBSx0043) was identified in *S.agalactiae* <SEQ ID 139> which encodes the amino acid sequence <SEQ ID 140>. Analysis of this protein sequence reveals the following:

Possible site: 25

>>> Seems to have an uncleavable N-term signal seq

INTEGRAL Likelihood = -11.04 Transmembrane 20 - 36 (12 - 43)

----- Final Results -----

bacterial membrane --- Certainty=0.5416 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GF: CAB13378 GB: Z99111 ylbL [Bacillus subtilis]

Identities = 124/344 (36%), Positives = 199/344 (57%), Gaps = 21/344 (6%)

Query: 20 WIIGFAFLILVLASLVRLEPYLLEMPGGAYDIRSVLKVNKADKAGSYNFVAVSGQAT 79
W++ L+ VL+ ++LEPY+ FG A ++ S++KV + KGS + V V A
Sbjct: 9 WMLVILILVLASL--FKLPPYITTKGKATLASLKVGGYPR-KGSLSLATVKKVPPAN 65

Query: 80 PAQLVYAMLTPPEL-----SKKETTGGFNDYLRINQFYMETSQNESIQALKLANKQ 135
P ++A + P+ E+ S KKE G S+ Y++ N++SQ ++ A + A K+
Sbjct: 66 PFTYVWAKMHPYRIIVDESIRKE-----GSDKEYMKRQLQMKSSQENAVIAAYQAGKK 122

Query: 136 VSLTYKGVVVLNLAKNSTPKDLHLADTVTGWNGKSPFNSSQLIKYVAALHIGDKVKVQY 195
VS ++ G+Y ++ +N K ++ + D + +GK++++ +LI Y+++ GDKV ++
Sbjct: 123 VSYSPNGIYASSGVVENMPAKGKIEVDKILIGADGKNYQSAEKILIDYISSKAGDKVTLKI 182

Query: 196 TSQGKKKESVGVKVLKNGKNGIGIGLTDHT--VLSDVDPVDPNTEGVGQPSAGLMPTLA 253
+ K+K + + + + GIG++ +T+ V + +DF R +GGPSAGLN +L
Sbjct: 183 ERSEKIKRVTLTLAKQPFDEPRAGIGVSLYTDENVKVEPDIDPELIENTGGPSAGLMSLE 242

Query: 254 IYDQLKQKEDLXGRKLQAGTGTITBQNHGVGDIGGAGLVVSAAKGMDIPFVNNPIDKXA 313
IY+QL K D KG IAGTGTI+ +G VG IGG KVV+A K G DIFP FN N
Sbjct: 243 IYNLTKPDETKGYDIAGTGTITDIDVGVKPGPVGIDQKVVAAKAGKIDIPFAPNQGASN- 301

Query: 314 KKGKTKVQINVOEAKAAKRLGTMKIVPVQMVQQAIDYLKKT 357
 ++Y+ A AK + + MKIVPV +Q AIDYL K K
 Sbjct: 302 -----SDYKNAVKIAKDIDENMKIVPVDTMQDAIDYLNK 337

- 5 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 141> which encodes the amino acid sequence <SEQ ID 142>. Analysis of this protein sequence reveals the following:

Possible site: 23
 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -10.24 Transmembrane 10 - 26 (6 - 34)
 10 ----- Final Results -----
 bacterial membrane --- Certainty=0.5097(Affixmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

15 The protein has homology with the following sequences in the databases:

>GP:CAR13378 GB:Z99111 ylbL [Bacillus subtilis]
 Identities = 132/348 (37%), Positives = 198/348 (55%), Gaps = 16/348 (4%)
 20 Query: 1 MKRLKKIKWMLVGLALALISLLALFFPLPYIEMFGGAYDIRTLVQNGKEDKRKAGYQF 60
 M R K M L W + L L I + L F L P Y Y I D G A + + + + + V G + K G +
 Sbjct: 1 MLRKKHPSWMLV-LILILAVLS--PIKLPIYITKGEATELASLIVEGGYPS-KGSLGL 56
 25 Query: 61 VANGISRASLAQLLYAMLTFFTEISTAEITG-GYSDADFLRINQFYMETSQNAIYQAL 119
 + V + A + + A + P + E I E G S D + + + + M + S Q A + A
 Sbjct: 57 MIVKVGPNPFTYVWAKMHPYIIVPDESIEKEGESDKEYMKRQLQMKSSQENAVIAAY 116
 30 Query: 120 SLAGKPVTLIDYKGVVLDVNESTFKGTLHLADTVTVNGKQFTSSAELIDYVSHLKLGD 179
 AGK V + + G + Y V K G + D + + + G K + S + + L I D Y + S K G D
 Sbjct: 117 QKAGKGVSYFNGIYASVVENMPKANGKEVGDKIISADGKNYSARKLIDYISSKAGD 176
 35 Query: 180 EYTVQFTSDNKKPKGVGRIILKLN--GKNGIGIALTDHTSVNSDVTIPSTKGVGGSAG 237
 + V T + + + K K + + + + + G I G + L + V E + F + + G G S A G
 Sbjct: 177 KVTILKEEKEKKRVLTLTKQFPDPEPRAGISVSLYTDRIKVPSPDIDPEINIGGPSAG 236
 40 Query: 238 LMFTLDIYDQITKEDLRKGRITAGGTIGDKGEVGDIGGAGLKVAAARAGADIFFVPFN 297
 L M + L + I Y + Q + T K D K G I A G T G T I D G + V G I G G K V A A + A G D I F F P N
 Sbjct: 237 LMSLEIYNQLTKFDETKGYDLAGTGTIDVDGKVGFIIGDIDKVAAADKAGKIDFFAPQ 296
 45 Query: 298 PVDKEIKKVNPNNAISNYYEAKRAAKRLKTMKIVPVTVQEAALYLRK 345
 N + S + Y + A + A K + + M K I V P V T + Q + A + Y L K
 Sbjct: 297 -----NGASKSDYKNAVKIAKDIDENMKIVPVDTMQDAIDYLNK 335

An alignment of the GAS and GBS proteins is shown below:

Identities = 229/339 (67%), Positives = 276/339 (80%)
 45 Query: 17 LKWWIIGFAFLNLVLASVLRPLPYIEMFGGAYDIRSVLKNVKADKAKGSYNFVAVSVS 76
 + K W W + G L + + L L P Y Y I E M F G G A Y D I R + V L N K D K G + Y F V A V + S
 50 Sbjct: 7 LKWWLGLLALISLLALFFPLPYIEMFGGAYDIRTLVQNGKEDKRKAGYQFVAVS 66
 Query: 77 QATPAQVLYAMLTFFTELSKEETTGGPSNDYLRINQFYMETSQNAIYQALAKLAKQV 136
 + A + A Q + L Y A M L T F P T E + S + E T T G G + S + D + L R I N Q F Y M E T S Q N + I Y Q A L I A K V
 55 Sbjct: 67 RASLAQLLYAMLTFFTEISTAEITGGSADFLRINQFYMETSQNAIYQALSAGKQV 126
 Query: 137 SLTYKGVVYVNLAKNSTPKDRILHLADTVTVNGKSPKNSQLIKIYVAALHGLDKVKVQIT 196
 + L Y K G V Y L + + S T F K L H L A D T V T V N G K F + S + + L I Y V + L L G D + V Q A T
 60 Sbjct: 127 TLDYKGVVYVLDVNESTFKGTLHLADTVTVNGKQFTSSAELIDYVSHLKLDEVDVQPT 186
 Query: 197 SQGKKKSGVKVIGLKNKGIGIGLTDHTSVLSDVPDNTGEGVGSGLMFTLAID 256
 S K K + V G + + I K L N K N G I G I L D T E T V S + + P + T + G V G G P S A G L M F T L A I D
 65 Sbjct: 187 SDNKKPKGVGRILHLKNGKNGIGIALTDHTSVNSDVTIPSTKGVGGSAGLMTFLDIYD 246
 Query: 257 QLVKEDLRKGRKTAGTGTIEQNGHVGDIGGAGLKVSAAGKGMDFPFPNIDKNKAG 316
 Q + K E D L R K G R I A G T G T I E Q N G H V G D I G G A G L K V S A A G K G M D F P F P N I D K N A G K

Shict: 247 QITKEDLRKGRTIAGTGTIGKDGEVGDIGGAGLKVVAAAEAGADIFFVPNNPVDKEIKKV 306

Query: 317 KTKVQTNVQEAKAAAKRLGTKMKIVPVQNVQQAIDYLLK 355

+NY+EAK AAKRL TKMKIVPV VQ+A+ YL+K

Sbjct: 307 NPNAISNYEEAKRAAKRLKTKMKIVPVTTVQEALVYLRLK 345

A related GBS gene <SEQ ID 8479> and protein <SEQ ID 8480> were also identified. Analysis of this protein sequence reveals the following:

Linop: Possible site: -1 Crepd: 10

McG: Discrim Score: 8.26

GVH: Signal Score (-7.5): -4.04

Possible site: 25

```
>>> Seems to have an uncleavable N-term signal seq
```

ALOM program count: 1 value: -11.04 threshold: 0.0

INTEGRAL Likelihood =-11.04 Transmembrane 20 - 36 (12 - 43)

PERIPHERAL Likelihood = 4.51 70

modified ALOM score: 2.71

*** Reasoning Step: 3

----- Final Results -----

bacterial membrane --- Certainty=0.5416(Affirmative) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

GP|5531383| putative secreted protein {Streptomyces coelicolor A3(2)} Insert characterized
PIR|T36157|T36157 probable secreted protein - Streptomyces coelicolor Insert
characterized

ORF01344 (361 - 1362 of 1671)

GP|5531383|emb|CAB51015.1||AL096852(13 - 247 of 259) putative secreted protein
{Streptomyces coelicolor A3(2)} PIR|T36157|T36157 probable secreted protein - Streptomyces
coelicolor

 $\text{fMatch} = 7.1$

%Identity = 38.4 %Similarity = 57.6

Matches = 58 Mismatches = 61 Conservative Sub.s = 29

EKW**R*****V**K**N**RDPK**R**KHKS**L**G**L**LKW**I**TG**F**A**P**LLLV**L**S**A**IV**K**LPY**L**E**M**PGG**A**D**I**R**S**V**L**KV**N**KADKA**R**G**S**YNFV-----
| : : : : | :: : || : | :
ML**S**R**I**R**T**PQ**F**LAVCG**L**PVALLAT**A**LFA**P**L**P**FSVAQG**L**TADV-----
 10 20 30 40

924 954 984 1002
-KKKESVGKVGIKLNGSKNGIKTGIGLTDHTEVL-----DVVP
 :| |:: ||
-----LGKNRCAEVITISGAPHTASQGLRMVTTIA---KESQSATTAALFYLRMDKGVDVV
 50 60 70 130 140

[illegible]

1272 1302 1332 1362 1392 1422 1452 1482
KGKTKVQYNYGSAKAARIGTGKMKIVPVNQVQAIDLYLKKTKTQRVRASARLCFATFDYQSAMIV*QSL*EYYI*N
 | | | | | | | | | | | | | | | | |
-----AECSDAQAELEKGLRLIPVTTLGGAVDSLKALESSKGVDPAC
 220 230 240 250

SEQ ID 8480 (GBS39) was expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 12 (lane 9; MW 65.2kDa) and Figure 15 (lane 3; MW 40kDa).

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 45

A DNA sequence (GBSx0044) was identified in *S.galactiae* <SEQ ID 143> which encodes the amino acid sequence <SEQ ID 144>. This protein is predicted to be UDP-sugar hydrolase. Analysis of this protein sequence reveals the following:

```
Possible site: 17
>>> Seems to have no N-terminal signal sequence
----- Final Results -----
bacterial cytoplasm --- Certainty=0.3908(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
```

The protein has homology with the following sequences in the GENPEPT database:

```
>GP:CB35227 GB:Z99120 similar to hypothetical proteins [Bacillus subtilis]
Identities = 114/280 (40%), Positives = 173/280 (61%), Gaps = 9/280 (3%)

Query: 1 MTELIRILHLNDLHSHFENFPKVKRPFH---DNCAQPIETISLDGLDNIDKSHPLTEAS 56
M E +R+ H NDLHSHFEN+PK+ + +Q+ ET+ D+Gd+D+ +TEA+
Sbjct: 1 MKEKRLHHTNDLHSHFENFPKIVDYIEQKKEHQSDGEETLVFDIGHLDRPQFVTEAT 60

Query: 57 SGKANVQLANELGIELATIGRNEGVGLSKKOLDQVYKSDPTFVIVGNLKD-NIIEPSWAK 115
KXANV L+N L I+ A IONNEG+ L +L +Y +F F VIV NL D N PSWA
Sbjct: 61 FGKANVDLLARLHIDGAIGNNEGITLPHZELAAALYDHAEPFVIVSNLFDKNGRPSWAV 120

Query: 116 FYIIVETQQQTCLAPLAYTFPYKYTYEPNGWTFIDPDLCKLCHLQINIK-EANCRLIMS 174
FY I + G +AFL T PYY Y+ GWT+ D + +K I E+K +A+ +LS
Sbjct: 121 FPHKSLKNGMSIAPLGVTVPYFPYVYKLGWVTDALSIK--ETILEVKGQADIIVLIS 178

Query: 175 HLGIRFDTRIAQEPSEIDLIGANTHHLFEBGELINGTYLAAAGKYGRFVGSIDITFONH 234
HLGI D +A+ EID+I+ +RTHHL E+G++NG LA+A KYG +VG ++IT D+
Sbjct: 179 HLGILDQVAEAVPEIDVILESHHLLLEDQGVVNGVLLASAKKYGRYVGCVEITVDS- 237

Query: 235 TLKIDILISTCDTKQLTGVPSDSDLKRLSKQKVENLKKV 274
+ I T + + + +S + + + B+K+
Sbjct: 238 VQRINSKTSASVQNAEWGESAETKAFINEKERBAERKL 277
```

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 46

A DNA sequence (GBSx0045) was identified in *S.galactiae* <SEQ ID 145> which encodes the amino acid sequence <SEQ ID 146>. This protein is predicted to be UDP-sugar hydrolase. Analysis of this protein sequence reveals the following:

```
Possible site: 44
>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood = -0.48 Transmembrane 5 - 21 ( 5 - 21)
----- Final Results -----
bacterial membrane --- Certainty=0.1192(Affirmative) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>
```


A related GBS nucleic acid sequence <SEQ ID 960> which encodes amino acid sequence <SEQ ID 960> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

5 >GP: CAB15227 GB: Z99120 similar to hypothetical proteins [Bacillus subtilis]
Identities = 29/137 (21%), Positives = 71/137 (51%), Gaps = 13/137 (9%)

Query: 3 AMLFYAGADVAINSGLIIVCFEKD-FSRKNLHESLPHQMLAKLTVSSQELLEIYETIY 61
A+ + D+++NSG+I+ P + ++ +LH PH + + ++ +EL E ++

10 Sbjct: 305 ALKEWCETDISMVNSGVILGPLKAGFVTKLLDLRIRICPHINFVAVRLTGERLKETI--VH 362

Query: 62 CQQQFLAQOKIHGNGFRSKCFGEVLHSGFDYKN-----GKIVYNEKIDIAKEEVI 111
+ + Q +I G+GFRG+ G++++G + + +I N +DI+ ++

15 Sbjct: 363 AASBQMEQLRIKGLGFRGEVNGMVYAGVEVETKRLDDGITHVTRITLNGEDISKHKQYS 422

Query: 112 LVIVLQYFYASYFECLEK 128
+ ++D + F ++

Sbjct: 423 VAVLDMFTLGLKLFELIR 439

20 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 47

25 A DNA sequence (GBSx0046) was identified in *S.galactiae* <SEQ ID 147> which encodes the amino acid sequence <SEQ ID 148>. This protein is predicted to be unnamed protein product. Analysis of this protein sequence reveals the following:

Possible site: 29

30 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3567 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
35 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein differs from AX026665 at the C-terminus:

Query: 181 SAKGHFVIRKK 191
SAKCH + +K
40 Sbjct: 181 SAKCHLLFVRK 191

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 149> which encodes the amino acid sequence <SEQ ID 150>. Analysis of this protein sequence reveals the following:

Possible site: 37

45 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3974 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
50 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 110/205 (53%), Positives = 147/205 (71%), Gaps = 15/205 (7%)

Query: 1 MRKEVTPPEMLNKNYK+GGPQIHFENIVKSDDIKFPQLVINKKSAFDVTVGQRFSEILLKY 60
 M+KE++PEM NYNK+GGP+FLHFE VK++ I+ L+ + K+AFD T FQGR++E+LLKY
 Sbjct: 9 MKKELSPENMYNKNF+GGPKFIHFEEQVKGADILLKLDVKNADTTTSFGQRYTFEVLKY 68

5 Query: 61 DFIVGDWGNQQLRLRGFYKDIATIRKNSRISRLDYIKETCNFGCAYFVLNPNFPROIKF 120
 D+IVGDWGNQQLRL+GPTKD+ I+K +RISRLDYIKE+CNFGCAYFVLN +P+DIKF
 Sbjct: 69 DYIVGDWGNQQLRLRGFYKSDDIKKTNRISRLDYIKETCNFGCAYFVLNLMHPQDIKF 128

10 Query: 121 DDERPHIKRKS-----RSKSQSSKSQTRNNRSQSNA-----NARFTSKKRKDTKRR 166
 ++ER +R+KS R K S Q +S+S N FTS+KR+ +
 Sbjct: 129 EBERQPRRKSPKSKSNRKRPNYSNQAPATPKSKSKRASKEQFENQAFTSQKRKSNTHK 188

Query: 167 QERHIKESODKEMTSAKQHFVIRKK 191
 +E+ K Q ++ + HF+IRKK
 15 Sbjct: 189 KEKS-KRNQTSQLNTKISHFIIRKK 212

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 48

- 20 A DNA sequence (GBSx0047) was identified in *S. galactiae* <SEQ ID 151> which encodes the amino acid sequence <SEQ ID 152>. Analysis of this protein sequence reveals the following:

Possible site: 32

>>> Seems to have no N-terminal signal sequence

25 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3627 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

30

A related GBS nucleic acid sequence <SEQ ID 9607> which encodes amino acid sequence <SEQ ID 9608> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

35 >GP:BA05225 GB:AP001515 unknown conserved protein [Bacillus halodurans]
 Identities = 205/349 (58%), Positives = 258/349 (73%), Gaps = 5/349 (1%)

Query: 18 PSIYSLTRDELIAWAIEHGEKKFRASQIWDWLYKKRVQSFDEMINISDPIALLNENFVV 77
 PSIY+L +EL W E GE KFRA+QI++WLY+KRV+ F EMIN+SID A L ++F +
 Sbjct: 17 PSIYTLQFEELWMLKQCEPKFRATQIFEWLYKRVQSFDEMINISDLRAKLEKHFLN 76

40 Query: 78 NPLKQRIQVQESADOTVKYLPDPQMLIETVLMRQHYGLSVCVITVQVGNIGCTPCASGL 137
 LK Q+S+DGT+K+LPEL DG IETVNR +YG SVCVITVQVC +GCTPCAS L
 Sbjct: 77 TLLKTVIQQSSDGTIKFLRLHDGYSIETVMRHNYSVCVITVQVQRLGCTPCASGL 136

45 Query: 138 IKKQDLNANGEITAQIMLVQKYFDERGQGERVSHIVVMGIGEPFDNYTNVLKFLRTVND 197
 +R+L GEI AQ++ Q+ DS QGERV IVMGIGEPFDNY ++ FL+TVN D
 Sbjct: 137 GGLKRNLEAGEIVAQVVEQQRANDS--QGERVSIVMGIGEPFDNYQALMFLKTVND 194

50 Query: 198 NGLAIGARHITVSTGLAHKIREFANEGVQVNLAVSLHAPNDLRSSIRINRSPFLEKL 257
 GL IGARHITVSTSG+ KI +FA+EG+Q+M A+SLHAPN +LRS +M +NR++EL KL
 Sbjct: 195 KGLNIGARHITVSTSGVVPKVIYQFADGELQINFAISLEAPNTELRKLMFVNFAHLEKL 254

Query: 258 FAALIEYIETINRIVTFEYIMINGVNDTPENAGELADLTCKIRKLSYVNLIPYNPVSSEH 317
 AI YVI+ T RRVTFEY + G ND E+A+ELADL K I+ +VNLIP NVE D
 55 Sbjct: 255 MDAIRYIDIKTRRRVTFEYGLGGRNDQVEHAEELADLKD I--CHVNLIPYRVSPERD 312

Query: 318 QYSRSPKERVEAFYDVLKNGVNCVVQBEHGTIDAAACQQLRSNTMKRD 366
 Y R+P++++ AF LK+ GVN +R+E G DIDAACQQLR+ K +
 Sbjct: 313 -YVTFPRDQIFAFERTLKRGVNVTIRREQGHIDIAACQQLRAKERKEE 360

60

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 153> which encodes the amino acid sequence <SEQ ID 154>. Analysis of this protein sequence reveals the following:

Possible site: 17

5 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

10 bacterial cytoplasm --- Certainty=0.2320 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 316/353 (89%), Positives = 339/353 (95%)

15 Query: 17 KPSIYSLTRDELIAWAIEHGEKKFRASQIWDNLYKKRVQSFDMNTISKDFIALINENFV 76
KPSIYSLTRDELIAWA+G G+K+PRA+QIWDNLYKKRVQSF+EMNTISKDF++LN++F
Sbjct: 2 KPSIYSLTRDELIAWAVERGQKQFRATQIWDNLYKKRVQSFEMNTISKDFVSIINDSFC 61

20 Query: 77 VNPLKQRIVQESADGTVKYLFELPDGKLIETVLMRQHYGLSVCVTTQVGCNIGCTFCASG 136
VNPLKQRIVQESADGTVKYLFELPDGKLIETVLMRQHYG SVCVTTQVGCNIGCTFCASG
Sbjct: 62 VNPLKQRIVQESADGTVKYLFELPDGKLIETVLMRQHYGHSVCVTTQVGCNIGCTFCASG 121

Query: 137 LIKKQRDLNNGEITTAQIMLVQKYFDERGGQGRVSHVVMGIGEPFDNYTNVLKFLRTVND 196
LIKKQRDLN+GEITTAQIMLVQKYFD+R QGERVSH+VVMGIGEPFDNY NV+ FLR +ND
25 Sbjct: 122 LIKKQRDLNNGEITTAQIMLVQKYFDDRKQGRVSHVVMGIGEPFDNYKNVMCFRLVND 181

Query: 197 DNGLAIGARHITVSTSGLAHKIRFANEGVQVNLAVSLHAPINDLRSSIMRINRSFPLEK 256
DNGLAIGARHITVSTSGLAHKIR+FANEGVQVNLAVSLHAPINDLRSSIMR+NRSFPLEK
30 Sbjct: 182 DNGLAIGARHITVSTSGLAHKIRDFANEGVQVNLAVSLHAPINDLRSSIMRVNRSFPLEK 241

Query: 257 LFAAIEYYIETNRRVTFEYIMLNQVNDTPENAGLADLTGKIRKLSYVNLIPYNPVSSEH 316
LF+ALEYYIE TNRRVTFEYIMLN VND+ + AQELADLTG IRKLSYVNLIPYNPVSSEH
35 Sbjct: 242 LFSAIEYYIETNRRVTFEYIMLNQVNDTIKQAGLADLTGKIRKLSYVNLIPYNPVSSEH 301

Query: 317 DCYRSRSPKERVEAFYDVLKKNQVNCVVRQEHGTDIDAACQQLRSNTMKDRQK 369
DCYRSRSPKERV AFYDVLKKNQVNCVVRQEHGTDIDAACQQLRS TMK+DR+K
35 Sbjct: 302 DCYRSRSPKERVAFYDVLKKNQVNCVVRQEHGTDIDAACQQLRSKMKDKREK 354

40 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 49

A DNA sequence (GBSx0048) was identified in *S.galactiae* <SEQ ID 155> which encodes the amino acid sequence <SEQ ID 156>. This protein is predicted to be VanZF. Analysis of this protein sequence reveals the following:

45 Possible site: 47

>>> Seems to have an uncleavable N-term signal seq

50 INTEGRAL Likelihood = -9.61 Transmembrane 86 - 102 (77 - 106)
INTEGRAL Likelihood = -8.80 Transmembrane 19 - 35 (15 - 42)
INTEGRAL Likelihood = -5.15 Transmembrane 113 - 129 (109 - 134)

----- Final Results -----

* bacterial membrane --- Certainty=0.4843 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
55 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAF36806 GB:AF155139 VanZF [Paenibacillus popilliae]

-103-

Identities = 45/154 (29%), Positives = 68/154 (43%), Gaps = 36/154 (23%)

Query: 17 RRFVQMLVILYCLIVRMCFGPQIMIEGVSTPNVQRFGRIVAL-----LVFPNSPESL 69
 R F+K+ V ++ L +V M G NV GR L L+P+S
 5 Sbjct: 36 RHLFVWYVFLFALVYMGTG-----IGNVWVGRYETLRVSEINLLPFS--- 82

Query: 70 DQLTSFKRIFWVQGVVVILLPLIIGLLSLKPSLRKYSVILLAFIMSIPIRQTVV 129
 + +T++ ++NI+L PL L ++ P R K+ F S+ IE TQ++
 10 Sbjct: 83 EGVTTY-----ILNILLMPLGFLPTWQPRTIKNTACTGFFFSIAIEALQL 132

Query: 130 LDILIDANRFVFIIDMLNITLGGPFAWYRNK 163
 +R+ IDDL NTLG YR K
 Sbjct: 133 -----NRRITDIDDLNITLGAIGYLLYRAFK 160

15 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 50

A DNA sequence (GBSx0049) was identified in *S.galactiae* <SEQ ID 157> which encodes the amino acid sequence <SEQ ID 158>. This protein is predicted to be multidrug resistance-like ATP-binding protein mdl. Analysis of this protein sequence reveals the following:

Possible site: 30

>>> Seems to have no N-terminal signal sequence

25	INTEGRAL	Likelihood = -6.79	Transmembrane	18 - 34 (17 - 36)
	INTEGRAL	Likelihood = -5.15	Transmembrane	247 - 263 (242 - 268)
	INTEGRAL	Likelihood = -2.81	Transmembrane	160 - 176 (158 - 176)
	INTEGRAL	Likelihood = -2.71	Transmembrane	141 - 157 (134 - 158)
	INTEGRAL	Likelihood = -1.12	Transmembrane	56 - 72 (56 - 73)
30	INTEGRAL	Likelihood = -0.69	Transmembrane	278 - 294 (277 - 294)

----- Final Results -----

bacterial membrane --- Certainty=0.3718 (Affixative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 35 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:BA805055 ABC transporter (ATP-binding protein) [Bacillus halodurans]
 Identities = 284/575 (49%), Positives = 406/575 (70%), Gaps = 2/575 (0%)

40 Query: 1 MSIIKLNWFFKEKKRYLIGILSLVAVNLIPPKINGSVIDAITGKLRPQLLWNL 60
 M + +LWFFK+KKK Y GI+ L++V++L L+PP++G ++D I G LT P LL +
 Sbjct: 1 MKVFDLWFFKQKKYSVGPQIVNLAIIVELLIVPEPRVVGIIIDVHYEOTLMPVLLQWI 60

45 Query: 61 LQLVLSALAMYGLRYIWMYILGTSYKLQVWRYRLFEHFTMSPFSFYQKRTGDIAMHA 120
 L AL +Y RY+WR+ I G S +L +++R +L+ HFT M+ FYQ+RTGDIAMHA
 Sbjct: 61 GVLAALALIVYARYLMKVMIPGASLRARLENQLYTHFTMMAFFYQKHRTGDIAMHA 120

50 Query: 121 TNDINSLRLAAGGVMSAVDSITALVTLITMPTTISQMILIAVILPLMALATSKLGR 180
 TNDI ++ AG GV++ VD+ ++TM TISW++TLI+++P+PLNAL TS G
 Sbjct: 121 TNDIRAIQTAGQGVLTLDVSLTMGGFVILTMATITISMKLTLSLPLMALITSYGS 180

Query: 181 KTHETFKESQAFFSELNNKQVSVSGVKVTSFGYQEQRIASFQEVNMTFVKNRMITY 240
 H+ F +QAFS LN+KVQSV+GV+VTK+FG +EQ+I +F++ + KN+
 55 Sbjct: 181 LHKRPHRAQAFSSINDKQVSVTVGVTVTAFOQRQDIAFRKQSDVVKNVAVARV 240

Query: 241 DVMFDPVLVIFIGASYVILAMGAFNISQGVTVGDVLTFTVYLDMLVPLMAGFLFM 300
 D +FDP +L +G SY L + GA + Q+TG L +F YL +L+WP++A GFLEH+
 Sbjct: 241 DALFDPPTISLIVGLSFLAIVFGARFVIEQLTIGQLTSPTLYGLLLINMIAFGFLNI 300

60

Query: 301 VQGGSVSYNRINSLLEQESDITDPLNFRPVVNGTLRYDIDFFRYDN--ESTLADIHPTL 358
 V+RG SYNRR++ LL++ +ITD I G+ ID FYN E LAD+ FL
 Sbjct: 301 VERGRASYNVSLQLQAKQETDSRARIHVPTGHDVAIDQFVYPNQKEPALAVQFEL 360

5 Query: 359 EKGQTLGLVGSQTSGKTSLIKLLLRHHVDVQGGKITLNKHDIRDYRLSELRLQLIGVVPQDQ 418
 +G+TLG+VG+TG+GKT+L++LL RR+D+ QG I L+ I Y L L+ G VPQD
 Sbjct: 361 SEGETTGLGVGKTGAGKTTLLRLQLQREYDQQGTIILDGRPIGHTYLDALKAAPGTVPQDH 420

10 Query: 419 FLFATSILENVRPNPTLSINAVKATKLAVHYDDIKOMPAGFETLIGEGVSLGGGQKQ 478
 FLF+ +I +M+ F P +I+ + +LAH++DDI Q G++T++GH+GV+LSGGGQKQ
 Sbjct: 421 FLFSATLADNIAFAKPDATISELIQVLSQLAHDDIIOFQGYDVTVGERGVTLGGGQKQ 480

Query: 479 RIAMSRAMILDPPDILLDDSLSAVDAKTEHAIIENLKNRQKSTTISAHRLSAVVHADL 538
 R++++RA++ +P+ILLDDSLSAVDAKTE AI+ +L+ R+GK+TII+AHRLGA+ HAD
 15 Sbjct: 481 RVSIRALLANPWLILLDDSLSAVDAKTEAILSSLAERKCKTITITAHRLSAIKHADH 540

Query: 539 ILVMQDGRVIERGQHQLLNKGGVYAFYASQOLE 573
 ILVM DGR++ERG H+ L+ GGWY Y QOLE
 Sbjct: 541 ILVMDGRIVERGTHETLMEAGGYYRNMVYRQOLE 575

20

There is also homology to SEQ ID 8.

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 159> which encodes the amino acid sequence <SEQ ID 160>. Analysis of this protein sequence reveals the following:

Possible site: 23

25 >>> Seems to have an uncleavable N-term signal seq

INTEGRAL	Likelihood	Transmembrane	176 - 192 (173 - 197)
INTEGRAL	Likelihood = -4.78	Transmembrane	267 - 283 (265 - 285)
INTEGRAL	Likelihood = -4.09	Transmembrane	18 - 34 (15 - 40)
INTEGRAL	Likelihood = -2.13	Transmembrane	151 - 167 (150 - 169)
INTEGRAL	Likelihood = -0.69	Transmembrane	85 - 101 (85 - 101)

30

----- Final Results -----

bacterial membrane	--- Certainty=0.4100 (Affirmative) < succ>
bacterial outside	--- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm	--- Certainty=0.0000 (Not Clear) < succ>

35

An alignment of the GAS and GBS proteins is shown below:

Identities = 172/609 (28%), Positives = 315/609 (51%), Gaps = 58/609 (9%)

40 Query: 1 MSIIKILWFFKFKKRYLIGILSLSAVLNLIIPPKNIGSVDAITTKGLTRPQLMLNL 60
 M + W++FK + + + +L L + P + G + + GK+ + + +
 Sbjct: 2 MKTARFNFYFYRYRFSFTVIAVAVLATYLVQKAPVFLGSELTL--GKIGQAYYVAM 59

45 Query: 61 LGLV----LSAL--AMGLRYIWRMYILGT---SYLKGQV-----RYRLFETHTOM 103
 G LSA M+ L + +L S+ L +VV R LF ++
 Sbjct: 60 SQGTHFSDPLSAPNAVMPKILMTYPTFLVLANLYLSFLTRVVSHTNRMRKGLFKLERL 119

50 Query: 104 SPSFYQKYRTGDLMAHATINDINSLTRAGQGVMSAVDASITAVLTILMTFTTISQM-- 160
 + +F+ +++ G++++ T+D+++ + +++ S+ +VT I++ + W M
 Sbjct: 120 TVAFFDRHKDGELGRFTSDLDN-----IQNSLNGSLIQVITIALYIGLVMVFRQ 171

55 Query: 161 -----TLIAVIPLPLMALATS-KLGRKTHETFKESQANPSLNKNQVESGVGVKTEKF 213
 IA P+ L+ L+ +L RK Q S LN + E+SG K
 Sbjct: 172 DSRLLATLTIASFVVALIFLWNIIRLAKYNTI--QQCEVSALNAPMDTISGQKAIIVQ 228

Query: 214 GYQEQLASF---QEVNQMTFVKIMRT-----MTYDVMFDPLVLLFGASVLT-LAM 262
 G OE + +F + V Q TF + + + M + + + +F+G++ VL++ +M
 Sbjct: 229 GVQEDWTATFLGHNRRVQATFKRLRSQQLFPVNMGMSLINTALVITPVGSTIVLSKDKM 288

60 Query: 263 GAFMISKGQVTVGDLVTFVYILDMLWPLMAIGFLNMVVGSGSVSYNRINSLLEQESDIT 322
 A +G +VTFV Y P+M I + +Q +RI + ++ ++
 Sbjct: 289 PA-----AAALGLVTFVYQYSQQYQPMQIASSWGELQLAFTGAHRIQEMFDETEVR 342

Query: 323 DPLNPLRPVVMGTLRYD-IDFFRYDNEETLADIHFTEKQQLGLGQVGSSEKTSILKLL 381
 P + + + +DF ++ L+D+ KG+ + +WG TGSSEK++ L+
 Sbjct: 343 PQNAFAF+SLKKEVAINHVDVFGVLPGLQKVLSDVSIAPKGMUAVGVPGSGKTIIMNLI 402

5 Query: 382 LREHDVTGQKITLNKHDIRDYRLSELRLQGLYVPOQQLFATSLLENVRFGNPTLSINAV 441
 R +DV G IT + DIRDY L LRQ +G V Q+ LF+ +I +N+RFG+ T+S + V
 Sbjct: 403 NRPYDWDAGSITIPKGRDIRDYDIADSLRQKGVILQKSVLPSGTITITONIRFDQYTSIQMNV 462

10 Query: 442 KKATKLARHVYDIDKQMPAGFETLIGKGVSLSGGQKQRIAMSRAMILDPDILILDDLSA 501
 + A + H++D I +P G+ T + + S GQKQ I++R ++ DP++LILD++ S
 Sbjct: 463 ETAARATHIDFIMSLPKGYNTYVSDDDNVFSTGQKQLISARTLLIDPEVILDEATSN 522

Query: 502 VDAKTHALLENKLNKQKSTLIIASHRLSAVVHADLLIWMQDQGVIERGQGHQLIMKGG 561
 VD TE I ++ G+++ +AHRLL +++AD I+V++DG+VIE+G H EIL++ G
 15 Sbjct: 523 VDTVTSKIQRAMEAIVAGRTSFVIAHRLKTLINADHIIVLADGKGVIERGNNHELLEKQG 582

Query: 562 WYAEITYASQ 570
 +YAE Y +Q
 20 Sbjct: 583 FYAELHMQ 591

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 51

A DNA sequence (GBS0050) was identified in *S. agalactiae* <SEQ ID 161> which encodes the amino acid sequence <SEQ ID 162>. This protein is predicted to be mdIB (ATP-binding prot). Analysis of this protein sequence reveals the following:

Possible site: 39

30 >>> Seems to have no N-terminal signal sequence

INTEGRAL	Likelihood = -8.65	Transmembrane	164 - 180 (155 - 183)
INTEGRAL	Likelihood = -5.15	Transmembrane	25 - 41 (21 - 46)
INTEGRAL	Likelihood = -4.88	Transmembrane	143 - 159 (133 - 163)
INTEGRAL	Likelihood = -1.49	Transmembrane	251 - 267 (251 - 270)
INTEGRAL	Likelihood = -1.33	Transmembrane	61 - 77 (61 - 77)

35 ----- Final Results -----

bacterial membrane	---	Certainty=0.4461 (Affirmative)	< succ>
bacterial outside	---	Certainty=0.0000 (Not Clear)	< succ>
bacterial cytoplasm	---	Certainty=0.0000 (Not Clear)	< succ>

40

The protein has homology with the following sequences in the GENPEPT database:

>GP:BA06054 ABC transporter (ATP-binding protein) [Bacillus halodurans]
 Identities = 278/582 (47%), Positives = 398/582 (67%), Gaps = 6/582 (1%)

45 Query: 1 MMKSNQWQVFKGLISYLRPYKWTPLALSLLLLTIVVKNIIPLDASHFIDHYLT-NVKNQ 59
 + Q VFKRL+SY YK ++A LL+ T + + P+I FID YLT T
 Sbjct: 9 LSSKQRTVYFKRLSYAAHYKGLMVAFLLPLATGAQLGPIIVKIFIDYLTFRPYPT 68

50 Query: 60 AVLILVG--YYSMYVLQTLIQYFGNLFARVSYISIVDIRRDANMERLGMYSFDRTEPA 117
 VL L+G Y +++ +I Y+ F +V+ SIV+ +R D F++++RLG+SFD+TEPA
 Sbjct: 69 DVLFLGAGYVLVHLTAVIDYQLFLQKVALSIVQRLRIDVPSVQRLGLSFFDQTEPA 128

Query: 118 GSIVSRITNDTEALSMDPGLSSSFISAIFITVITYMLMDIKITGLVALLLPVILF 177
 G +VSRITNDTE+I +++ +L++F+ I M- L++ L +LLP+IF L
 55 Sbjct: 129 GGLVSRITNDTESIKELYVTIVLATFVQNIILPLIGFAAMPYLVNLTALYILVLLPLIFAL 188

Query: 178 VNVYRKKSFTVIACTRSLLSDINKSEISIRIGIRIVQAFQGEERLCTEPEINKEHVVA 237
 + VYRK S A LS +N +++ESI+G+ I+Q F QB R++ EF IN EH +
 Sbjct: 189 MQVYRKYSRFFYADMSKLSLNGRINESIQMAIIQMFRGRMKKPSAINDRHFLAG 248

60 Query: 238 NRSMDLSLFLRPAGSLLLKLAYAVLMAYFQFTGVKGLTAGLMYAFIQVYNRLFDLIE 297

```

+SM LD L LRPA+ +L +LA ++++YFG + + G++YAF+ Y+R F+P+ +
Sbjct: 249 MKSMKGLGILLRPAVVDLILALMLLSYFGIMGMOTAVEIGVVYAFVNYLDRFPFVFNQ 308

5 Query: 298 VTONFSTILOTSMVSAGRVFDLIDETGFSQKNT--AFVREGNIEFKMVSFSDYGGKQI 355
+ S .Q ++VSAGRV L+D P ++ E A + BGN+EF+MVSPSYDYGK +
Sbjct: 309 MMRGLSMFQQAIVSAGRVFKLMDHRKLAPDRGNEHPAIIIGBNVEFRMVSPSYDYGKTVN 368

Query: 356 LBNVSPVSKKGETLAFVGATGGGKSIINVFMRFYEPFGQGVLLDGDKIDRVYSQEQLRN 415
L N+SF+VKKGET+H VG TGGK+SIINV MRFY Q G++L+DGK + + +LR
10 Sbjct: 369 LBNISFTVKKGETVALVGHGTGGKTSINVIMRFYPLDGGILIDGKPLTSFENNELRAK 428

Query: 416 IGLVLQDPFLYHGTIKSNIMY-ODITDOEVODAAEFVDADQFIQKLPKDYDAVRSEGS 474
+GLVLQDPFLY GTI SNI++Y Q I+D ++ AA FV AD FI++L Y+ V+ERG+
15 Sbjct: 429 VGLVLQDPFLYGTITAGNIRLYDQAI SDORIKRAASFVRADGFIERLSHGVTYKVTERRA 488

Query: 475 SFSTGQROLIAFARTVASKPKILLDEANTANDSETEQIVODSLAKMRQRTTIIAIAHRL 534
+FS+GQROLL+PART+ +P ILILDEATA++D+ETE+ +Q++L +M+QRTTIIAIAHRL
Sbjct: 489 TFSQORQLLSFARTMVREPAILLDENTASVDTEATQEALEMRKQRTTIIAIAHRL 548

20 Query: 535 STIODANCITYVLDRKIIESGNHSLDLKGLTYRMVLOAG 576
STI+DA+ I VL +G+I+ E G R+ L+ KG Y +MY LQ G
Sbjct: 549 STIKDADQILVLHQGEIVERGTHDELIAKGLYKMYVLQKG 590

```

There is also homology to SEQ ID 160.

- 25 A related GBS gene <SEQ ID 8481> and protein <SEQ ID 8482> were also identified. Analysis of this protein sequence reveals the following:

```

Lipop: Possible site: -1 Crend: 10
McG: Discrim Score: -4.63
GVH: Signal Score (-7.5): -5.85
30 Possible site: 39
>>> Seems to have no N-terminal signal sequence
ALOM program count: 5 value: -8.65 threshold: 0.0
INTEGRAL Likelihood = -8.65 Transmembrane 164 - 180 ( 155 - 183)
INTEGRAL Likelihood = -5.15 Transmembrane 25 - 41 ( 21 - 46)
35 INTEGRAL Likelihood = -4.88 Transmembrane 143 - 159 ( 133 - 163)
INTEGRAL Likelihood = -1.49 Transmembrane 251 - 267 ( 251 - 270)
INTEGRAL Likelihood = -1.33 Transmembrane 61 - 77 ( 61 - 77)
PERIPHERAL Likelihood = 3.02 483
modified ALOM score: 2.23
40 *** Reasoning Step: 3
----- Final Results -----
bacterial membrane --- Certainty=0.4461(Affirmative) < succ>
45 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the databases:

```

ORF01277(322 - 2028 of 2340)
50 BGAD|108578|BS0971(2 - 667 of 673) hypothetical protein [Bacillus subtilis] OXNH|NIGLBS1137
conserved hypothetical protein GP|2226165|emb|CA74449.1||Y14080 hypothetical protein
[Bacillus subtilis] GP|2633307|emb|CA312811.1||Z99109 similar to ABC transporter (ATP-
binding protein) [Bacillus subtilis] PIR|H69828|H69828 ABC transporter (ATP-binding
55 protein) homolog ybeH - Bacillus subtilis
%Match = 28.5
%Identity = 40.8 %Similarity = 69.1
Matches = 234 Mismatches = 171 Conservative Sub.s = 162

162 192 222 252 282 312 342 372
60 RLPLQHIHQYLLCYOTLS*LCKTAESSESVSIKSC*IRVVGMLKRMPSN*KNRKHLMKSN*QWVFKELISYLRPYKWF
: : : : : : :
MKIGKTLNRYALLVKKLL
10

```

[illegible]

There is also homology to SEO IDs 330, 4634 and 5788.

50 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 52

A DNA sequence (GBSx0051) was identified in *S.agalactiae* <SEQ ID 163> which encodes the amino acid sequence <SEQ ID 164>. Analysis of this protein sequence reveals the following:

55 Possible site: 25

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

```
60      bacterial cytoplasm --- Certainty=0.0635(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>
```


A related GBS nucleic acid sequence <SEQ ID 9609> which encodes amino acid sequence <SEQ ID 9610> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```

5  >GP:AAA25224 GB:M07483 anthranilate synthase beta subunit
    [Lactococcus lactis]
    Identities = 101/191 (52%), Positives = 133/191 (68%), Gaps = 4/191 (2%)

10 Query: 14 MLLLVNDYDSFTYNLQKYLVSVEVFVTKNDVNLFLLAESAEIAVLSPGPGHPKDSAGM 73
    M+L+DNDYDSFTYNL QY+ V +V V+KND +L +AE A+A++ SPQPG P DAGM
    Sbjct: 1 MLLIINDYDSFTYNLQYVGVLTDAVAVKNDSDSLGWAEKADALIFSPGPGWEDAGM 60

    Query: 74 VELINQFIGKKPILGICLGHQALAECLGRLANLHVHMGKQSVVTINDHTSLFEGIDSP 133
    LI QF G+KPIILGICLG QA+ E GG+L LA+ VMHGK S V +F +S
    Sbjct: 61 ETLLIQFAGQKPIILGICLGPQAIQVFGGLRLARQVMHGKNSOVROTSGLNLIPIHLEPK 120

    Query: 134 TQVMRYHSLVVTDD---LPENIAVIARSNEDEIMAFHCPSLKVYAMQFHPESIGSIDGMK 190
    VMRYHS+V+ + LP+ A+ A+ +D EIMA ++Y +QFHPESIG+DGM
    Sbjct: 121 FLVMRYHSIVMDEVALEP-FAITAVATDDGEIMATENEKEQIYGLQFHPESIGTLOGMT 179

20 Query: 191 MIENFLTLI 201
    MIENF+ +N+
    Sbjct: 180 MIENFVNCVNE 190
  
```

25 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 165> which encodes the amino acid sequence <SEQ ID 166>. Analysis of this protein sequence reveals the following:

```

    Possible site: 57

30 >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
            bacterial cytoplasm --- Certainty=0.3183 (Affirmative) < succ>
            bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
            bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
  
```

An alignment of the GAS and GBS proteins is shown below:

```

    Identities = 104/186 (55%), Positives = 131/186 (69%)

40 Query: 14 MLLLVNDYDSFTYNLQKYLVSVEVFVTKNDVNLFLLAESAEIAVLSPGPGHPKDSAGM 73
    M+LL+DNDYDSFTYNL QYLS + E V+ N ENL+ +A+ A A+VLSPPGP PK+A +M
    Sbjct: 1 MLLIINDYDSFTYNLQYLSSEFDEIVTLVNDQENLYDMAKKNALVLSPPGMPKKNQM 60

    Query: 74 VELINQFIGKKPILGICLGHQALAECLGRLANLHVHMGKQSVVTINDHTSLFEGIDSP 133
    +LI F KPIILG+CLGHQA+AE LGS L LA VMHG+QS + SLF+ +
    Sbjct: 61 PKLIQDFYQTKPIILGICLGHQAIAETLGGTILRLAKRVHMGROSTIETQGPASLFRSLPQS 120

    Query: 134 TQVMRYHSLVVTDLPEINIAVIARSNEDEIMAFHCPSLKVYAMQFHPESIGSIDGMKIE 193
    VMRYHS+VV LP+ +V AR +D RIMAF +L ++ +QFHPESIG+ DGM MI
    Sbjct: 121 ITVMRYHSIVVDLQKGSFVSRTARDCDDQIMAFEBHTLPLFLGLQFHPESIGTPOGMTMIA 180

50 Query: 194 NPLTLI 199
    NF+ I
    Sbjct: 181 NFIAAI 186
  
```

55 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 53

A DNA sequence (GBSx0052) was identified in *S.agalactiae* <SEQ ID 167> which encodes the amino acid sequence <SEQ ID 168>. Analysis of this protein sequence reveals the following:

Possible site: 58

```
>>> Seems to have a cleavable N-term signal seq.
INTEGRAL    Likelihood = -8.17    Transmembrane  117 - 133 ( 108 - 140)
INTEGRAL    Likelihood = -1.70    Transmembrane  150 - 166 ( 150 - 166)

----- Final Results -----
      bacterial membrane --- Certainty=0.4270 (Affirmative) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
```

The protein has homology with the following sequences in the GENPEPT database:

```
>GP: CAB12877 GB:Z99109 similar to biotin biosynthesis [Bacillus subtilis]
Identities = 70/168 (41%), Positives = 106/168 (62%)

Query: 8   YIALMVALLIVLGFIPGIFPGFIPVPIVLQNLGVMLAGALLSGRKFGLAVAIFFLLVAIG 67
      +IA+ AL+ VLGF+P + L F PVPI LQ LGVMLAG++L + FL+ +FLLVA G
Sbjct: 9   HIAITALMAVLGFMEFLFLGFTFPVPTTQTLGVMLAGSILRPKSAFLSLVFLLLVAFG 68

Query: 68  APFLFQGRSGSLVTLFGPTAGYLLTYPPAAFFIGLGLKVKTKLWQFLIWIPIGVLLID 127
      AP LFOGR G   PGF+AG+L+ YP A++ I L +++ + F +PG++ I
Sbjct: 69  APFLFQGRGGFVFGFSGFLIAYPLASMLISLAANRLRKVTVLELFPTHIVEFGIIFIY 128

Query: 128 ICGSIVLSFQTSLEPLTKSLFSNLIIFIGDTLKASICLIYRKFNKRLT 175
      + G V +F + L++ F +L ++PGD +KA++ + K L+
Sbjct: 129 LIGPIVQAFIMHIDLSQAFMSIATVPGDLIKAAVSAPTAIKITQALIS 176
```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 169> which encodes the amino acid sequence <SEQ ID 170>. Analysis of this protein sequence reveals the following:

Possible site: 51

```
>>> Seems to have an uncleavable N-term signal seq
INTEGRAL    Likelihood = -10.03   Transmembrane  113 - 129 ( 109 - 139)
INTEGRAL    Likelihood = -8.97    Transmembrane  55 - 71 ( 52 - 76)
INTEGRAL    Likelihood = -7.54    Transmembrane  10 - 26 ( 6 - 38)
INTEGRAL    Likelihood = -5.79    Transmembrane  86 - 102 ( 81 - 105)
INTEGRAL    Likelihood = -2.87    Transmembrane  33 - 49 ( 28 - 51)
INTEGRAL    Likelihood = -1.97    Transmembrane  150 - 166 ( 150 - 168)

----- Final Results -----
      bacterial membrane --- Certainty=0.5012 (Affirmative) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
```

An alignment of the GAS and GBS proteins is shown below:

```
Identities = 80/168 (47%), Positives = 108/168 (63%), Gaps = 1/168 (0%)

Query: 3   TRTTTYIALMVALLIVLGFIPGIFPGFIPVPIVLQNLGVMLAGALLSGRKFGLAVAIFFLL 62
      T+   +A+M L+I+LGFIP IPLGFIPVPIVLQNLGVMLAG +LG +KG L+V +F L
Sbjct: 4   TKELVKVAMTTLIIILIGFIPATIPGLFIPVPIVLQNLGVMLAGLMLGKKGTLSVFLF-L 62

Query: 63  LVAIGAPFLFQGRSGSLVTLFGPTAGYLLTYPPAAFFIGLGLKVKTKLWQFLIWIPIFG 122
      ++ + P G R+ + L GP+AGY++ Y L + + + FL + I G
Sbjct: 63  VIGLFLFVFGSGRRTTIPVLMGSGAGYVIAYILVPIVPSLLYRNWFSKSTPLAFIALLISG 122

Query: 123 VLLIDICGSIVLSFQTSLEPLTKSLFSNLIIFIGDTLKASICLIYRKFI 170
      V+L+D+ G+I LS T + L SL SNL+FIQD+KA I II K+
Sbjct: 123 VVLVDVLGAIWLSAYTQMSLVTSLSLNVLPFGDTIKAIATIIAVKY 170
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 54

- 5 A DNA sequence (GBSx0053) was identified in *S. galactiae* <SEQ ID 171> which encodes the amino acid sequence <SEQ ID 172>. Analysis of this protein sequence reveals the following:

Possible site: 17
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3914 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S. pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

20 Example 55

A DNA sequence (GBSx0054) was identified in *S. galactiae* <SEQ ID 173> which encodes the amino acid sequence <SEQ ID 174>. Analysis of this protein sequence reveals the following:

Possible site: 15
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1864 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 9611> which encodes amino acid sequence <SEQ ID 9612> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

- 35 >GP:BAB05467 GB:AP001513 biotin synthase [Bacillus halodurans]
 Identities = 133/316 (42%), Positives = 201/316 (63%), Gaps = 2/316 (0%)
- Query: 17 NYIHLEDEILSGKTSISYEQALRIILNS-DENWWEIYAALYLKQVSRNIRNLNVLISAK 75
 N+I LA E++ GK IS +AL ILNS D+ + A ++ ++LN++++AK
 40 Sbjct: 2 NWIQLAQEVIEGKR-ISENEALAIILNSPDELLILLQGAFTIRQTYGKRVKIAMIMNAK 60
- Query: 76 QGLCAENCGYCSQSKESTADIDKFGLLPQNVILKQATVAHQNGASVPCIAMSGTYKSKRE 135
 G C ENCGYCSQS S A ID + ++ + IL+ A AH+ +CI SG P+ R+
 45 Sbjct: 61 SGPCFENCGYCSQSSISKAPIDAYPMVNEKETTLEGAKRAHELNVGTYCYVASGRGPTNRD 120
- Query: 136 IEQLQCVIPRIKISLFLSICITAGFLDRQELHLQKAGIDRINENLNATPERNNYNIATTH 195
 I+ + ++ RIK + L+IC G L EQ QLK AG+DR NHH+NT ++ I T+H
 50 Sbjct: 121 IDHVTAEVREIKDYGLKICACGLILKPBQAEQLKAGVDRYNNHNVTSARHSDQITTS 180
- Query: 196 SFKDRCDTLERIHEDIVCSGFICGMGESDEGLITLAPRLKELDYPYSIPVNFLLAVGDT 255
 +++DR +T+E + + I CSG I GM E+ E ++ +AF+L+ELD SIPVNF L A++GT
 Sbjct: 181 TYEDRVNTEVVKHSGISPCGVI VGMKRTKEDVVMNAFQLRELDADSI PVNPLHADTGT 240

-111-

Query: 256 PLGKYNLYLTPKCLKIMAMLR FVFFPKRLSNGREVHFNFSVLTLVVDSTFLGNYLT 315
 PL + LTPI CLK+++ R+V P KE+R+S GREV+ ++ + L +S F+G+YLT
 Sbjct: 241 PLQGVHELTPIYCLKVLSELYVCPTKRIISGGREVNLSLQPLGLYAANSIFIGDYLT 300

5 Query: 316 EGGRNQHTDIEFLKEL 331
 G+ + D + L+ L
 Sbjct: 301 TAGQRETADHQILKDL 316

No corresponding DNA sequence was identified in *S.pyogenes*.

- 10 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 56

A DNA sequence (GBSx0055) was identified in *S.galactiae* <SEQ ID 175> which encodes the amino acid sequence <SEQ ID 176>. Analysis of this protein sequence reveals the following:

- 15 Possible site: 24
- >>> Seems to have no N-terminal signal sequence
- 20 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3440(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

- 25 A related GBS nucleic acid sequence <SEQ ID 9613> which encodes amino acid sequence <SEQ ID 9614> was also identified.

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

30 Example 57

A DNA sequence (GBSx0056) was identified in *S.galactiae* <SEQ ID 177> which encodes the amino acid sequence <SEQ ID 178>. Analysis of this protein sequence reveals the following:

- Possible site: 15
- 35 >>> Seems to have no N-terminal signal sequence
- Final Results -----
 bacterial cytoplasm --- Certainty=0.1985(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 40 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

- 45 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 58

A DNA sequence (GBSx0057) was identified in *S.galactiae* <SEQ ID 179> which encodes the amino acid sequence <SEQ ID 180>. Analysis of this protein sequence reveals the following:

Possible site: 32

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.11 Transmembrane 347 - 363 (347 - 363)

----- Final Results -----

bacterial membrane --- Certainty=0.1044 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CA11722 GB:AL445064 acetyl-CoA acetyltransferase related
 protein [Thermoplasma acidophilum]
 Identities = 113/388 (29%), Positives = 181/388 (46%), Gaps = 31/388 (7%)

Query: 4 RDVYIGFGLRTPIGIKGQFKHYR-PELLGAHLNQIKKIESSESNID----SIICQNTV 57
 RDV+I RT IG G+ F + P+L GA IK + E+++D +I GN +
 Sbjct: 2 RDVFIAAKRTAIGKFGRSFSKIKAPQLQA----AIKAVMDRAHVDPASVEEIVIMZVI 57

Query: 58 --GTGGNIGRLINTLFSDYESIYPVQTIDMQCSSSSALFFGYLKISTGINKVLVGGIES 115
 G G N + + T+++ CAS A+ +I+ G + V+ GG+ES
 Sbjct: 58 QAGINGQNFAGQAAFPHGGLPNSVLKTYTVNVVCSAGMLAVESAAREIALGERDLVIAGGMS 117

Query: 116 SSLQPMR-----RYAKEDNRNGEYTVAQ-PSDPSYAEIVMLE----GAQRVQKYGFERRE 165
 S P R+ + Y + D + E A+R +K+G RE
 Sbjct: 118 MSNAPFLLPADLRNGPKHLHLHONYKIDDMITDGLLDATYFERHMGVSAERTSRKFGITRE 177

Query: 166 MLDKLAFLSHKRALTAQGGYLEEVILPWEGM-RDQGVRLKETFFQKLEPIMENSPILT 224
 M D+ + S++R+ A + G + I+ EG+ D+G+RK +LP + + +LT
 Sbjct: 178 MADEYSVQSVGERALRATESGEFADSIYQFEGLDHDSGIRKATIMEDLARLEPAFDKNGILT 237

Query: 225 IGVNCLMHDAAPFLTLQSQRT--BFRIVHIVEVAG-----DPKLSPELVHTEATEKLLTE 276
 GN + D + L + S+K E+ + I + G DP E AT KLL +
 Sbjct: 238 AGNSAQLSDGGSSALMIASEKAINETGLKFIARTITGYEQRLDPLDFVEAPIPATRKLLBK 297

Query: 277 THTKISDYDAIENNEFFAAIDALFNHYYPEEREKFNIPGGTLAYGHFPAACSGIINIHLIM 336
 H I YD +E NE F+ + + + E+FN+ GG +A GHP SG I+ LM
 Sbjct: 298 QHKISDYIDLVEHNEAFSIAIVTVRNELKIDNERFNVNGGVAIGHPIGNSGARIIIVTLM 357

Query: 337 QALKYKRNKMLGTLAAGAGGVGMAISIE 364

ALK+++ GL + GG +++E
 Sbjct: 358 NALKIRHLKTGLATLCHGGGAHTITLE 385

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 181> which encodes the amino acid sequence <SEQ ID 182>. Analysis of this protein sequence reveals the following:

Possible site: 22

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -1.28 Transmembrane 345 - 361 (345 - 361)

----- Final Results -----

bacterial membrane --- Certainty=0.1510 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP:BA03328 GB:AB035449 acetyl-CoA c-acetyltransferase
 [Staphylococcus aureus]
 Identities = 115/382 (30%), Positives = 184/382 (48%), Gaps = 29/382 (7%)

Query: 1 MIDVYIAAGLRFTPIGLVGHQFAKEQPEILGAKLINALQNKYFV---PIDQVIGCNTVGTG 57
 M I A R T G G +PE L L + KYP ID V+ GN VG G
 Sbjct: 1 MNQAVIVAAKRTAPGKYGTGLKHLEPEQLLKPILFQHPKPKYFEVTSKIDVDVLGNVVG 60

Query: 58 GNIGRLMTLYSHLGEVSALTVDQMCASGAALSGVYAKIKAGMASNLVGGIESBS--- 114
 GNI R L + L +S+ +T+D Q C S ++ I+AG + GG+ES+S
 Sbjct: 61 GNIAKALLLEAGLKDSPGVITDRCQSGSGLSEVYACRMICAGAGKVYLAGVVESTERP 120

Query: 115 ---LQPEGVYAGADWRQGAAYKVAQSPDSGISPFAMIEGAERVAREHGPTKEYLNHWTLRS 171
 +P SVY +A Y+ A F+P+ P +MI+GAE VA+ ++E + + RS
 Sbjct: 121 WKIKRPHSGVYETA--LPEPYERASFAPEKEDP-SMIQGAENVAQYDVSRELQDEFAYRS 177

Query: 172 HQKASYCQBQALLADLILDLSGA-----SDQGIPELSSKVLKVPPTLGBGHVISAANA 226
 HQ + + ++ IL ++ +D+ ++ + + P++ +G ++RAN+
 Sbjct: 178 HQLTAENVKIGNISQREILITVKGSEIFNIDESLKSHIPKDNFGRKPVII-KGTVTAAS 236

Query: 227 CLTHDAAPFLQLSSQPSAFL-----IDVVEVAGDPQRSPLMVIKASQVLEKHGLG 278
 C+ +D A L+ + +A+LL D V V D + + A LL++ L
 Sbjct: 237 CMKNDGAVLLIMKIMAYELGFEHGLLFKDGVTGVGVDSNFPICGIPALSNLKRQQT 296

Query: 279 MADMTAIEWNEAFVADIGLFETHYPDLLDRYNI PGALAGHPHYGASAAIILHLMRALE 338
 ++ IE NEAF+ + +NI+GGALA GHYPYAS A ++ L +
 Sbjct: 297 IENTIEVIEINEAFSAQVQACQALNININIQNLNIWGGALAGHPYASGAQLVTRFLYMD 356

Query: 339 IKKRGYGAIAAAGGQGFVL 360
 + IA++ GG G A L
 Sbjct: 357 KET---MIASMGIGGLGNAAL 375

30 An alignment of the GAS and GBS proteins is shown below:

Identities = 182/362 (50%), Positives = 243/362 (66%), Gaps = 2/362 (0%)

Query: 5 DVYIGPLRTPIGLVGHQFAKEQPEILGAKLINALQNKYFV---PIDQVIGCNTVGTG 64
 DVYI GLRTPIG+ GKQF +PE+LGA L+N ++ + ID +IGCNTVGTG 64
 Sbjct: 3 DVYIAAGLRFTPIGLVGHQFAKEQPEILGAKLINALQNKYFV---PIDQVIGCNTVGTG 61

Query: 65 RLMTLYSHLGEVSALTVDQMCASGAALSGVYAKIKAGMASNLVGGIESBSLQPSVY 124
 RLMT+ S + T+DQMCAS+ +AL GY KI G+ +LWGISBSLQPS Y
 Sbjct: 62 RLMTLYSHLGEVSALTVDQMCASGAALSGVYAKIKAGMASNLVGGIESBSLQPSVY 121

Query: 125 AKEDNRNGEYTVAGFSPDSYARTVMEBGAQRVCQKYGPPEMLDGLAPLSHKRALTAKG 184
 A D R G Y VAQSPDSG + M+EGA+RV ++GF +E L+ SH+A+ ++
 Sbjct: 122 ASADWRQGAAYKVAQSPDSGISPFAMIEGAERVAREHGPTKEYLNHWTLRS HQKASYCQBQ 181

Query: 185 GYLSEVILPMEGMRDQGRV-KLKETFPQKLPRIMENSPLLTIGNVCLMHDAAFLTLQGG 243
 L ++IL+ G DQ+R +L K+P ++ +++ N CL HDAAFL TL QGG
 Sbjct: 182 ALLADLILDLSGASDQGIPELSSKVLKVPPTLGBGHVISAANA CLTHDAAPFLQLSSQ 241

Query: 244 KTEPRIVHIVEVAGDPKLSPELVHTATEKLLTETHFKISDYDAIKENRPFADIDLPHY 303
 + F+++ +VEVAGDP+ SP +V A++ LL+ ++D AIEKNE FA ID LF +
 Sbjct: 242 PSAPFLIDVVEVAGDPQRSPLMVIKASQVLEKHGLGADMTAIEWNEAFVADIGLFETH 301

Query: 304 YPEERKFNIPGGTFLAYGHYPYCSGIINILHMQALKYKNKPMGLFALAGAGGVGMALIGBY 365
 YP+ +++NIPGG LAYGHYPY S I ILHM+A+L KN G+ AIA AGG G A+ ++Y
 Sbjct: 302 YPDLLDRYNIPGALAYGHYPYASAAIILHLMRALEIKNGRYGIAAIAAGGQGFVALLKY 363

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 59

60 A DNA sequence (GBSx0058) was identified in *S. agalactiae* <SEQ ID 183> which encodes the amino acid sequence <SEQ ID 184>. Analysis of this protein sequence reveals the following:

Possible site: 13

- >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -3.82 Transmembrane 149 - 165 (149 - 165)
- 5 ----- Final Results -----
 bacterial membrane --- Certainty=0.2529(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>
- 10 The protein has homology with the following sequences in the GENPEPT database:
 >GP:CAB12876 GB:Z99109 similar to long-chain fatty-acid-CoA ligase
 [Bacillus subtilis]
 Identities = 90/382 (23%), Positives = 158/382 (40%), Gaps = 24/382 (6%)
- 15 Query: 47 ISTHSLNQLVRFVSKLQKALPIICKWILTHNRISRLKEV--QYAPQLADFGVLSST 104
 IS L+ L F +KL P++ N +IS + P+ + +SG+
 Sbjct: 95 ISNADLVVTLAFFINKLTDSTFPVLLDNCA-DISEAADPLTIDPEHPFMGTSGS 153
- 20 Query: 105 TADAKLLNRSPTNSDFFSIQNAVFSVTSNSKLFIQGDFPTGNLNLALSLLLOGTIVV 164
 T K RS SW + F+ FS++S+ K+ I G + L A+S L LSGT+ +
 Sbjct: 154 TGKPKAFTRSHRSNWSSEFTCTETDFSISSDDKVLIPGALMSSEFLYGAVSTFLGGTVCL 213
- Query: 165 TQMSVKYKWTMEKTVTHLYLLPSYIKLVEQYSKETALDNKTIITSSQVSDSLLEG 224
 +K S + + ++ LY +P+ + + K I + + + +S + +
 Sbjct: 214 LKGPSPAKAKEWLCRESISVLYTVPTMTALARIEGFPDPSVKI ISSGADNPAS -KKKL 272
- 25 Query: 225 YRKEPKVSVKIPFGASELNVSVWDGDIRDKPQVGEIVPNVAVRKE----- 273
 P + + FYG SEL++V++ D + KP G NV + I+
 Sbjct: 273 AAANPFLKLYDFVTSSELSFVTFSSPEDSKRKPFSAGRPFNVRKIEIRNAGGERCPGEI 332
- 30 Query: 274 GRIFVKTPYSICG-----LSSEYCAQGYGELID--GKLYLFGRGGDQNCQGI KLYLPL 326
 G+IFVK+P G E+ D +D G LY+ GR G+ ++ +
 Sbjct: 333 GKIPVKSFMRFSGYGVNGSTPDENMTVDMMGYVDEEGFLYISGRINGMIVYGLNLPFERI 392
- 35 Query: 327 IEKIKTCPYKIDNAVFTKESQSHQSGESHCCVILBNQMCQCECLNLSHFEEKYVGFH 386
 + CP ++ A + G+ + V+ N + W + K +
 Sbjct: 393 ERVLLACPEVSEANVVGIPDEYNGEIA--VAVILGNANARTLKACCKQLASTYKIPKGV 450
- 40 Query: 387 IVSKIPLPMGGKIDYQKLKRQL 408
 +P SGKI ++K+ L
 Sbjct: 451 FADSLPETSSEGGIARSRVKKWL 472

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 185> which encodes the amino acid sequence <SEQ ID 186>. Analysis of this protein sequence reveals the following:

- 45 Possible site: 52
 >>> Seems to have no N-terminal signal sequence
- 50 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.2487(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

- 55 Identities = 154/413 (37%), Positives = 235/413 (56%), Gaps = 9/413 (2%)
- Query: 1 MLESIAKTVKRTNSDKLPGD-LQVSYGEFYNLR-QDMASQDNKRVISTHISLNLQVLR 58
 ML L+ K +KK D + ++Y E + V +D +D+ ++IS LNLQ+
 Sbjct: 1 MLTKLEYMAKQCFNKKAIVADQISLITYQLQAVLTKDTIKDSVPYITGISRYLNLQLS 60
- 60 Query: 59 FVSKLCQKALPIICKFNLT--ENEISRLKEKVVQAPQLADFGVLSSTADAKLLMRSF 115
 P+ L + + PII FN++ +I ++ R+ + ADF VLSSGTT AKI NR
 Sbjct: 61 FLRLGLKSGSCPTILIHFNISGTFQQQIKHVDGRL--KKADFVLSSTGKAKLFWRL 117

Query: 116 TSMDFSSIQNAYPSVTSNSKLFIQGDFSTGNNLALSLLLGGTLVVTQKMSVKVQT 175
 ++W+ F QN P +T NS LF+ G PSFTGNNLAL+ L GG LV++QK S+K W +
 Sbjct: 118 STWTRLEFDYQNKVFGWIGNSCLFLHGGSPSTGNNLALALQWAGGCLVLSQKLSKTLWS 177

5 Query: 176 LWRKTGVTHLYLLPSYKLVQYSKETALNNTIT+SSQYVSDDLGLYRKHPKVSVKI 235
 LW+ V+HLVLLP+YL + Y + + ++TSSQ +S Ld Y+K P++ + I
 Sbjct: 178 LMQAKKVSHULYPTIYNIILPYIITKNNWTALHLLTSSQMSQELLRHYYKKPQLKIVI 237

10 Query: 236 FYGASELNMYVMYDGRDIRDKPQYVGHIVENVAVRIKGRIPVKITPYSCIGLSSRYCAGD 295
 FYGASEL++W+GR VG+ P+V++ K+ IFV+TPYS+ G+S Y D
 Sbjct: 238 FYGASLSFTWNGRAAVKINGLAVGQPPFDVSIStKDKRIPIVETPSVSGMSQYTSVSD 297

15 Query: 296 YGELJGKLYLPGRGWQCNQSGIKLYLPRLEKIKTCPIYKDAVAPTKRQSQHQSSHC 355
 G++ L L GR DW NQ G+K +L P L+E F +K+A A K + +
 Sbjct: 298 LGMSPAGLILBGRQDQWVWQRGVKKHLESLVELAHQAQFNVKBAHL- KIGKGBETLIL 356

Query: 356 CIVLIENQMCQECMLWSEHFEKKYGFKNHYHVSkipLMPGKIDYQQLKROL 408
 +VL + +L+ + K+Y ++ +PL +GKI+ + L ++
 Sbjct: 357 VLVLTKKDCIAPIKDFIALYLNBSGLPKYLYIDCLPLKNGKINREVLNKKI 409

20

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 60

A DNA sequence (GBSx0059) was identified in *S. galactiae* <SEQ ID 187> which encodes the amino acid sequence <SEQ ID 188>. This protein is predicted to be endonuclease III (pdg). Analysis of this protein sequence reveals the following:

Possible site: 46

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.00 Transmembrane 25 - 41 (25 - 41)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.1001(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

30

35

The protein has homology with the following sequences in the GENPEPT database:

>GP:BAH05417 GB:AP001512 endonuclease III (DNA repair) (Bacillus halodurans)
 Identities = 95/202 (47%), Positives = 134/202 (66%)

40 Query: 1 MLKAKSRYIIRIILKLPFDKPSIDFTNVELLVAVMLSCATTDAAVNVKTPALFERFP 60
 ML+K +++ + I ++PDA+ L +N FELL+AV+LSAQ TDA VNKVTP LF ++
 Sbjct: 1 MLTKKQTQEAVALADIMYFDACELTYSNPFELLIAVVLSCATDAAVNVKTPALFAKYK 60

45 Query: 61 NPLVLAQADPKIEPIYSKIGLYRNKARPLMQCAQLIEHFDGKVPRTIRQELSLAGVGR 120
 P +E+E I IGLYRNKA+ + + L+E + G+VP+ R EL LAGVGR
 Sbjct: 61 TPEDYIAVPLEELEQDIRSIGLYRNKAKNIIKIQSLIEQYGGVEFQDRDLVVKLAGVGR 120

50 Query: 121 KIANVVMVSFGGIPAZAVDVIHTRICGHQICQKQASPLIEKRVMEVPEEWLAHQ 180
 KIANVV SV FG+PA AVDTHV R+ K IC+ + ++E+ +M+ +P +EW +H
 Sbjct: 121 KIANVVASVAFGPEALAVDTHVERVSKRLGICRWKDNVTQVEQTLMKKIPNDWESISHER 180

Query: 181 MIYFGRAICHKNPKCDQYFQL 202
 +I+PGR C +NP+CD P L
 Sbjct: 181 LIFFGRYHCKAQNPQDCICPLL 202

55

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 189> which encodes the amino acid sequence <SEQ ID 190>. Analysis of this protein sequence reveals the following:

Possible site: 44

>>> Seems to have a cleavable N-term signal seq.

----- Final Results -----

- 5 bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

- 10 Identities = 91/199 (45%), Positives = 133/199 (66%)
- Query: 2 LSKAKSRVYIRRIIKLPDAKPSLDFTNVFELLVAVMLSAQTDDAANKVTFALPFRFPN 61
 + KA+ ++ I ++FP+AK LD+ F+LL+AV+LSAQTTD AVNKVTP L++ +P
 Sbjct: 3 IGKARLAKVITIGQMFPEAKSELOWRTFPQLLIHAVLSAQTTDKAVNKVTFGLMQSYPE 62
- 15 Query: 62 PWLAQADPKIEPTYSKIGLYRNKARFANCAQLBHFDPKGVPTQRLESAGVGRK 121
 LA A+ ++E + IGLY+NKA+ + + A+ + + F G+VP+T +ELES L GVGRK
 Sbjct: 63 IRDLAFAELSDVENALRTIGLYKNKAKNIKTAQAIRDDFKGQVPKTHKELSLPGVGRK 122
- 20 Query: 122 TANVVMVSFGGIPAFVDTHTVTRICKHHQICKQSAPLEIEKRVMEVLPPEMLAAHOSM 181
 TANVV++ +G+PA AVDTHV R+ K I A +IE +M +P ++W+ H +
 Sbjct: 123 TANVVLAEVYGVPALAVDTHTVARVSKRLNISSFDADVKQTEADLMAKIPKCDMIITHRL 182
- 25 Query: 182 IYFGRALCHPNPKCDQYP 200
 I+PGR C K PKC+ P
 Sbjct: 183 IFFGRYECIAKPKCEICP 201

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

30 Example 61

A DNA sequence (GBSx0060) was identified in *S. galactiae* <SEQ ID 191> which encodes the amino acid sequence <SEQ ID 192>. Analysis of this protein sequence reveals the following:

Possible site: 51

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

- 35 bacterial cytoplasm --- Certainty=0.2264 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 40 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:BAA96473 GB:AB036428 hypothetical 8.3 kDa protein [Streptococcus mutans]
 Identities = 53/67 (79%), Positives = 62/67 (92%)

- 45 Query: 1 MKVLPDVQNLKKGPIYVYIGKRLYDIEVMKIELQRLYDNGLSRDYVLAELILRREHR 60
 MK L+DVQ LLK+PGI+VY+GKRLYDIE+MKIEL+RLYDNGLS+ DY L AELILRREHR
 Sbjct: 1 MTLVLDVQRLKQGPVYVYIGKRLYDIEVMKIELQRLYDNGLSKSDYVLAELILRREHR 60
- 50 Query: 61 LELKLEN 67
 +E E+EN
 Sbjct: 61 IEKEREN 67

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 193> which encodes the amino acid sequence <SEQ ID 194>. Analysis of this protein sequence reveals the following:

Possible site: 57

>>> Seems to have no N-terminal signal sequence

-117-

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1962(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

5

An alignment of the GAS and GBS proteins is shown below:

Identities = 53/66 (80%), Positives = 60/66 (90%)

Query: 1 MKVLFDVQNLKKPGITVYVYLGKRLYDIEVMKRLQRLYDNKLSRDDYLKASLILRRHR 60
 MK L-DVQ LLK PGI-VV-GKRLYDIE+MKRLQRLYD+GL+ + DYL ASLILRRHR
 Sbjct: 7 MKTLYDVQQLKMPGIFVYLGKRLYDIE+MKRLQRLYDGLDRLDYLNASLILRRHR 66

Query: 61 LELEKE 66
 LELEKE

15 Sbjct: 67 LELEKE 72

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 62

20 A DNA sequence (GBSx0061) was identified in *S.agalactiae* <SEQ ID 195> which encodes the amino acid sequence <SEQ ID 196>. Analysis of this protein sequence reveals the following:

Possible site: 31

>>> Seems to have no N-terminal signal sequence
 25 INTEGRAL Likelihood = -0.06 Transmembrane 133 - 149 (133 - 150)

----- Final Results -----

bacterial membrane --- Certainty=0.1022(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 30 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GF:BA05144 GB:AP001512 glucose kinase [Bacillus halodurans]
 Identities = 145/315 (46%), Positives = 209/315 (66%), Gaps = 2/315 (0%)
 35 Query: 6 LGIDLGOTTIKFGILTEGEVQEKWAEINTLENGRHIVSDIVESLKHRLSYGLTKDDF 65
 +G+D+GOTTIK LF GE+ +N I TN + G I ++I ++L RLS + +K D
 Sbjct: 7 VGVDSGTTIKMAFLTAGEIVDKWEIPINKQDGGALITTNLADALDKRLSGHHSKSDL 66

40 Query: 66 LGIGGSGFGAVDRSTKVTGAFNLNMDTQFVGSGVIEKEVGIPFFINDANVAALGEHV 125
 +GIG+G+FG ++ + + A N+ W D + +E+E +P +DNDAN+AALGE W
 Sbjct: 67 IGIGGSGFGPIEMDTGPIYHAVNIGWRDPF-LKDKLEETKLPVIVINDANIAALGEMWK 125

45 Query: 126 GAGANNPDVVPVFTLGTGCGGQVIADGNLHGVAAGGRIGHMIVDPENGPTCTCGNKGL 185
 GAG +++ +TLGTGCGGG++A+GN++HGV G GEIGH+ V DE G C CG GCL
 Sbjct: 126 GAGDGAKNLLTTLGTGCGGIVANGNIIHGKMGAGETGHTTIVPEGAPCNCGKTGCL 185

50 Query: 186 ETVASATGVVKVARQIARQYBSSATKAINDSDITVTSKDIPIAHEIDDKFANSVVERVS 245
 ETVASATG+ R+A + + + + S + D +T+KD+P A+ D FA SVV+ ++
 Sbjct: 186 ETVASATG+IARATGVTSEK-RSQIALQYDKRGVLTAKDVPSAADAFAFALVVOHIA 244

55 Query: 246 RYLGAANANISNIIINPDSPVIGGVSAAECPLRSRVKRYPTFAFPQVKKSTKIKAEKG 305
 YLG A AN++N LNP+ +VIGGVS AG+ L ++++F +A P+V + +IA LG
 Sbjct: 245 YYLGAANLANANPEKIVIGGVSAGDYLKPKIHPFAYALPRVADGAERIALTG 304

Query: 306 NDAGIIGAASLANQQ 320
 NDAG+IG L QQ
 Sbjct: 305 NDAGVIGGGLVKKQ 319

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 197> which encodes the amino acid sequence <SEQ ID 198>. Analysis of this protein sequence reveals the following:

Possible site: 23

5 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

10 bacterial cytoplasm --- Certainty=0.1060 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 270/319 (84%), Positives = 292/319 (90%)

15 Query: 1 MSKILGLIDLGSTTIKFGILITLGEVQEKVAIETNTLENGRHIVSDIVESLKHRLSLYGL 60
MS+KLLGLIDLGSTTIKFGILT GEVQEKVAIETN LE G+HIV DI+ S+KHRL LYGL
Sbjct: 1 MSQKILGLIDLGSTTIKFGILITAAAGEVQEKVAIETNILEGGKHIVPDIASIKHRLDLYGL 60

20 Query: 61 TKDDFLGIGMNSPGAVDRSKTVTGAFNINWADTQEVGVSVEKEVGIPFFIDNDANVAAL 120
+ DF+GIGMNSPGAVDR + TVTGAFNINW +TQEVGSV+EKE+GIPF IDNDANVAAL
Sbjct: 61 SSADFVGIGMNSPGAVDRDINTVTGAFNINWKTQEVGVSVEKELGIPFPAIDNDANVAAL 120

25 Query: 121 GERWVGAGANNPDVVFVTLGTGVGGGVIADGNLIHVAGAGGEIGHMIVDPENGFTCTCG 180
GERWVGAG NNPVVF+TLGTGVGGG+IADGNLIHVAGAGGEIGHMIV+PENGFTCTCG
Sbjct: 121 GERWVGAGNNPDVVFMTLTGTGVGGGIIADGNLIHVAGAGGEIGHMIVEPENGFACTCG 180

30 Query: 181 NKOCLETVASATGVVVRARQLAEQYEGSSAIIKAIDNGDVTVSXDIPIAAEDGDKFANSV 240
+ GCLETVASATGVV+VAR LAE YEG SAIIKAIDNG+ VTSXDIPI+AAE GD FA+SV
Sbjct: 181 SHGLETVASATGVVVKARLLAAYEGSDSAIIKAIDNGOVSXDIPIAAEAGDGFADSV 240

35 Query: 241 VERVSYLLGLAANISNIIINPDSSVYIGGVSAAAGEFLRSRVEKYFVTFAPQVKKSTIK 300
VE+V YLGLA+ANISNIIINPDSSVYIGGVSAAAGEFLRSR+EKYFVTF FQV+ STIK
Sbjct: 241 VEKVGYYLGLAANISNIIINPDSSVYIGGVSAAAGEFLRSRIEKFVTFPFQVRYSTIK 300

35 Query: 301 IAEIENDAGIIGAASLANQ 319
IAEIENDAGIIGAASIA Q
Sbjct: 301 IAEIENDAGIIGAASLANQ 319

40 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 63

A DNA sequence (GBSx0062) was identified in *S.galactiae* <SEQ ID 199> which encodes the amino acid sequence <SEQ ID 200>. Analysis of this protein sequence reveals the following:

Possible site: 19

45 >>> Seems to have a cleavable N-term signal seq.

----- Final Results -----

50 bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

55 >GP:CA814385 GB:Z99116 similar to hypothetical proteins [Bacillus subtilis]
Identities = 51/124 (41%), Positives = 71/124 (57%), Gaps = 1/124 (0%)

Query: 3 MSVILLIIVILLAFVAVASNNYKVRRAAKFLDNESFQKIMSEGGQLIDISAGAFHKKIL 62
MS +++++I AF+ + +Y +R K L E F+ +QLID+RE F HTL
Sbjct: 1 MSNMVILIIIFPAFTIYMIASYVYQCRIMKTLTEEFPRAGYRKAQLIDVREPNFEGHIL 60

Query: 63 GARNIPASQFKVALSALRKDKPVLLYDASRGQSIFRIVLLLRKSGFNOLYVLKDGFNWYT 122
 GARNIP SQ K + +R DKPV LY + +S R LRK G + +Y LK GF W
 Sbjct: 61 GARNIPQLSKQRKRNIRTDKPVLYCQMSVRS-GRAAQTLRKNGCTEYLNKGGFKKGG 119

Query: 123 GRVK 126
 G++K
 Sbjct: 120 GKIK 123

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 201> which encodes the amino acid sequence <SEQ ID 202>. Analysis of this protein sequence reveals the following:

Possible site: 30
 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -4.41 Transmembrane 4 - 20 (1 - 22)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.2763(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>P:BAB06532 GB:AP001516 unknown conserved protein [Bacillus halodurans]
 Identities = 46/120 (38%), Positives = 64/120 (53%)
 Query: 8 LMLLVGIVGYTYWYFSPFRMAKQVDNETFKDVMRQQLIDLRPAAFRTKHILGARNF 67
 +WL+L+ ++ Y + K K + E F R+ QLID+REP + + HILGARN
 Sbjct: 5 VWLVLLALLVYVLKRLTYTEKYLKTLIQEFPQGYRKAQLIDVRREPREDYSGHILGARNI 64
 Query: 66 PAQQPDAAIKGLRKDKPVLIYENMRPQYRVPAVKIKKAGFEDVYVLKDGIDYMDGKVKQ 127
 P Q +K +R D+PV +Y + R A KK G EDV LK G W GK+K+
 Sbjct: 65 PLSQLKQRLKEVRTDQPVLYCQSGARSQAAILKKKHGVEDVNLHKGGRFQVTKGKIK 124

An alignment of the GAS and GBS proteins is shown below:

Identities = 63/126 (50%), Positives = 85/126 (67%)
 Query: 1 NMMSVILLIIVILLAFVMAQSMYWRVRAAKFLDNESFQKEMSRGQLIDIREAGAFHRKH 60
 M +++ ++L+ V + +WNY+ R+ AK +DNE+F+ M +GOLID+RE AF KH
 Sbjct: 1 MSPITILLMLLVGIVGYTYWYFSPFRMAKQVDNETFKDVMRQQLIDLRPAAFRTKH 60
 Query: 61 ILGARNIPASQFKVALSALRKDKPVLLYDASRGQSIFRIVLLLRKSGFNOLYVLKDGFNWYT 120
 ILGARN PA QF A+ LRKDKPVL+Y+ R Q V L+K GF +YVLKDG -Y
 Sbjct: 61 ILGARNFPAQQPDAAIKGLRKDKPVLIYENMRPQYRVPAVKIKKAGFEDVYVLKDGIDY 120
 Query: 121 WTRGVK 126
 W G+VK
 Sbjct: 121 WDGKVK 126

A related GBS gene <SEQ ID 8483> and protein <SEQ ID 8484> were also identified. Analysis of this protein sequence reveals the following:

Lipcp: Possible site: -1 Crend: 1
 McG: Discrim Score: 17.55
 GvH: Signal Score (-7.5): 3.36
 Possible site: 17
 >>> Seems to have a cleavable N-term signal seq.
 ALOM program count: 0 value: 8.86 threshold: 0.0
 PERIPHERAL Likelihood = 8.86 99
 modified ALOM score: -2.27
 *** Reasoning Step: 3
 ----- Final Results -----
 bacterial outside --- Certainty=0.3000(Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

5 >GP:CAE13350 GH:K99111 similar to GTP-binding elongation factor
[Bacillus subtilis]
Identities = 455/609 (74%), Positives = 534/609 (86%), Gaps = 2/609 (0%)

10 Query: 4 LRTDIRNVAIIAHVDHGKTTLVDELLEQSHTLDERKKLRKRAMDSNDLEKEGGITILAN 63
LR D RH AIIAHVDHGKTTLV D LL Q S T ++ ERAMDSND E ERGITILAN
Sbjct: 3 LRNDLRNIIAHVDHGKTTLV D QLLHQAGTFRANBQVARRAMDSNDLERGITILAN 62

Query: 64 TAVAYNDVRINIMTPGHADPGSEVERIMGMVGVVLAVDAYEGTMPQTRFVLKALEQN 123
TA Y D RINI+DTGHADPGSEVERIMGMVGVVLAVDAYEG MPQTRFVLKALEQN
15 Sbjct: 63 TAINYKDRINILDTGHADPGSEVERIMGMVGVVLAVDAYEGCMPQTRFVLKALEQN 122

Query: 124 LIPIVVVVKIDKPSARPEVDEVLELFELGADDDLPFVVYASAINGTSSMSDDPSD 183
L P VVVVKID+ ARP EV+DEV L+L FIEL A+++CL+FPVVYASAINGT+S+ DP
20 Sbjct: 123 LNPVVVVVKIDRPFARPEVIDEVL DLFELDANEQLEFPVVYASAINGTASL--DPKQ 180

Query: 184 QKTYAPIFOTIIDHIPAPVDNSEEPLOQVSLDYNDPVGRIIGRVFRGTVKVGQVT 243
Q+ M ++TII H+PAPVDN+EEPLQFQV+LLDYND+VGRIGIRVFRGT+KVG QV+
Sbjct: 181 QDENBEALYTTIIKHVPAPVDN+EEPLQFQV+LLDYNDYVGRIGIRVFRGTMKVQGV 240

25 Query: 244 LSKLDGTTKNFRVTKLPFFGLERKEIQEAGDILAVSGMEDIFVGETVPTFDALIEPL 303
L KLDGT K+FRVTK+PGF GL+R EI+EAAGDL+AVSGMEDI VGETV P D +PLP
Sbjct: 241 LMKLDGTAKSFRVTKIPGQGLKRVIEEAGAGDLVAVSGMEDINVGETVCPVDHQDPLP 300

30 Query: 304 VLRIDEPTLQMTFVNNSPFAGREGKNTSRKVERLLAEQLDVSLSRVDPDTSPOKMTV 363
VLRIDEPTLQMTF+VNNSPFAGREGK++T+R+K+ERL ++LQTVDSLRV+PT SPD W V
Sbjct: 301 VLRIDEPTLQMTFVNNSPFAGREGKVTARKIEERLQSQQLQTVDSLRV+PTSDPWV 360

Query: 364 SGRGELHLSILIEHMRREGYELQVSRPEVILKEIDGVQCEPFRVQIDPFEYQALIGS 423
SGRGELHLSILIE HMRREGYELQVS+PEVILKEIDGV+CEP ERVQID PEE+ G++++S
35 Sbjct: 361 SGRGELHLSILIEHMRREGYELQVSKPEVILKEIDGVRCPEFRVQIDPFEHTGSVMS 420

Query: 424 LSRKGDMLDMQVGNQGTSLIPLIPARGLIGVSTFSLMRGVGINNHTFDQYLPVQ 483
+ RKG+M+DM GNGQ RLIF +P+RGLIGVSTFSL+TRG+GI+NHTFD Y P+ G
40 Sbjct: 421 MGARKGEMVMINNGNQGVLIFTVP+RGLIGVSTFSL+TRGFGILNHTFDQYLPVQ 480

Query: 484 RIGGRHGRGLVSIENGKATYSIMRIERGTFIPNGIEVYEGMIVGNSRNDLGVNIT 543
++GGR +G LVS+ENGKAT+Y I IE+RG IFV PG EYVYEGMIVG++RNDL VN++
Sbjct: 481 QVGRRCQGLVLSMENGKATSYGIGIEDRGVIFVPGTEVYEGMIVGHNNDLGVNVS 540

45 Query: 544 TAKQNTNVSATKDQTAIVIKTFRILTLERSLFLADDEYMEVTPESIRLRQLINKARD 603
KQ TNVSATKDQ T IK RI++LESL+L +DEY EVTPESIRLR+ILNK R+
Sbjct: 541 KKKQNTNVSATKDQTTTIKARIMSLERSLEYLNDEYCEVTPESIRLRKLINKNER 600

50 Query: 604 KAKKKKSA 612
KA KKKK+A
Sbjct: 601 KAKKKKTA 609

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 205> which encodes the amino acid sequence <SEQ ID 206>. Analysis of this protein sequence reveals the following:

55 Possible site: 36
>>> Seems to have no N-terminal signal sequence

60 ----- Final Results -----
bacterial cytoplasm --- Certainty=0.1738 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 594/613 (96%), Positives = 607/613 (98%)

```

Query: 1  MTNLRTDIRNVAIIAHVDHGKTTIVDELLKQSHITLDERKGLERAMDSNDIEKRGITIL 60
      1  MTNLR DIRNVAIIAHVDHGKTTIVDELLKQSHITLDERKGLERAMDSNDIEKRGITIL
5  Sbjct: 1  MTNLRDIRNVAIIAHVDHGKTTIVDELLKQSHITLDERKGLERAMDSNDIEKRGITIL 60

Query: 61 AKNTAVAYNDVRINIMDTPGHADFGGEVERIMKMVGCVLVVDAYEGTINPQTRFVLKAL 120
      61 AKNTAVAYNDVRINIMDTPGHADFGGEVERIMKMVGCVLVVDAYEGTINPQTRFVLKAL
      61 AKNTAVAYNDVRINIMDTPGHADFGGEVERIMKMVGCVLVVDAYEGTINPQTRFVLKAL 120
10 Sbjct: 61 AKNTAVAYNDVRINIMDTPGHADFGGEVERIMKMVGCVLVVDAYEGTINPQTRFVLKAL 120

Query: 121 EQNLPIPVVNNKIDKPSARPSEVVDREVLSEFLRIGADDQDLPFPVYASAINGTSSMSDD 180
      121 EQNLPIPVVNNKIDKPSARPSEVVDREVLSEFLRIGADD+QLFPVYASAINGTSS+SDD
      121 EQNLPIPVVNNKIDKPSARPSEVVDREVLSEFLRIGADDQDLPFPVYASAINGTSSLSDD 180
15 Sbjct: 121 EQNLPIPVVNNKIDKPSARPSEVVDREVLSEFLRIGADDQDLPFPVYASAINGTSSLSDD 180

Query: 181 PSDQEKTMAPIDFTIIDHIPAPVDNSEPLQFQVSLDYNDVFGRIGIRVFRGTIVKVG 240
      181 PSDQEKTMAPIDFTIIDHIPAPVDNSEPLQFQVSLDYNDVFGRIGIRVFRGTIVKVG
      181 PSDQEKTMAPIDFTIIDHIPAPVDNSEPLQFQVSLDYNDVFGRIGIRVFRGTIVKVG 240
20 Sbjct: 181 PADQEHMAPIDFTIIDHIPAPVDNSEPLQFQVSLDYNDVFGRIGIRVFRGTIVKVG 240

Query: 241 QVTLSEKLDGTTNFRVTKLPGFGLERKIQEAKAGDLIIVSGMEDIPVGETIPTDIE 300
      241 QVTLSEKLDGTTNFRVTKLPGFGLER+RIQEAKAGDLIIVSGMEDIPVGET+TPTD +E
      241 QVTLSEKLDGTTNFRVTKLPGFGLERKIQEAKAGDLIIVSGMEDIPVGETIPTDIEVE 300
25 Sbjct: 241 QVTLSEKLDGTTNFRVTKLPGFGLERKIQEAKAGDLIIVSGMEDIPVGETIPTDIEVE 300

Query: 301 PLFVLRIDEPTLQMTFVFNNSPFAGREGKWTSRKVEERLLAELQTDVSLRVDPDTPDK 360
      301 PLFVLRIDEPTLQMTFVFNNSPFAGREGKWTSRKVEERLLAELQTDVSLRVDPDTPDK
      301 PLFVLRIDEPTLQMTFVFNNSPFAGREGKWTSRKVEERLLAELQTDVSLRVDPDTPDK 360
30 Sbjct: 301 ALPIILRIDEPTLQMTFVFNNSPFAGREGKWTSRKVEERLLAELQTDVSLRVDPDTPDK 360

Query: 361 WTVSGRGLHLSILIEYMRREGYELQVSRPEVIIKEIDGV+CEPPEFVQIDTPPEYQAI 420
      361 WTVSGRGLHLSILIEYMRREGYELQVSRPEVIIKEIDGV+CEPPEFVQIDTPPEYQAI
      361 WTVSGRGLHLSILIEYMRREGYELQVSRPEVIIKEIDGV+CEPPEFVQIDTPPEYQAI 420
35 Sbjct: 361 WTVSGRGLHLSILIEYMRREGYELQVSRPEVIIKEIDGV+CEPPEFVQIDTPPEYQAI 420

Query: 421 IQSLSERKGMELMDQMVGNQOTRLIPLIPLFARGLIGYSTFLSMTRGYGIMNHTPDQYLPV 480
      421 IQSLSERKGMELMDQMVGNQOTRLIPLIPLFARGLIGYSTFLSMTRGYGIMNHTPDQYLPV
      421 IQSLSERKGMELMDQMVGNQOTRLIPLIPLFARGLIGYSTFLSMTRGYGIMNHTPDQYLPV 480
40 Sbjct: 421 IQSLSERKGMELMDQMVGNQOTRLIPLIPLFARGLIGYSTFLSMTRGYGIMNHTPDQYLPV 480

Query: 481 VQGEIGGRHGRGALVSIENGKATTYSINRIEERGTIFVNPGLVYEGMIVGENSRDNDLGV 540
      481 VQGEIGGRHGRGALVSIENGKATTYSINRIEERGTIFVNPGLVYEGMIVGENSRDNDLGV
      481 VQGEIGGRHGRGALVSIENGKATTYSINRIEERGTIFVNPGLVYEGMIVGENSRDNDLGV 540
45 Sbjct: 481 VQGEIGGRHGRGALVSIENGKATTYSINRIEERGTIFVNPGLVYEGMIVGENSRDNDLGV 540

Query: 541 NITTAKQNTNVRSATQDQTAIVIKTPRIITLESLEFLADDEYMEVTPESIRLRKQLINKA 600
      541 NITTAKQNTNVRSATQDQTAIVIKTPRIITLESLEFLADDEYMEVTPESIRLRKQLINKA
      541 NITTAKQNTNVRSATQDQTAIVIKTPRIITLESLEFLADDEYMEVTPESIRLRKQLINKA 600
50 Sbjct: 541 NITTAKQNTNVRSATQDQTAIVIKTPRIITLESLEFLADDEYMEVTPESIRLRKQLINKA 600

Query: 601 ARDKANKKKKSAB 613
      601 ARDKANKKKKSAB
55 Sbjct: 601 ARDKANKKKKSAB 613

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 65

A DNA sequence (GBSx0065) was identified in *S. agalactiae* <SEQ ID 207> which encodes the amino acid sequence <SEQ ID 208>. This protein is predicted to be D-glutamic acid adding enzyme MurD (murD). Analysis of this protein sequence reveals the following:

RSD motif 441-443

Possible site: 29

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

```

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 9615> which encodes amino acid sequence <SEQ ID 9616> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```

5 >GP:AA095449 GB:AF068902 D-glutamic acid enzyme MurD [Streptococcus pneumoniae]
  Identities = 341/449 (75%), Positives = 394/449 (86%)

Query: 5 MKTITTFENKKVLVLGLARSGSAAARLLAKLGAIVTVNDGKPFDPNPTAQSLLLEGIKVV 64
  MK I P+NKKVLVLGLA+SGS+AAARLL KLGAIIVTVNDGKPF++NP AQ LLESGIKV+
10 Sbjct: 1 MKVIDQFQKKVLVLGLARSGSAAARLLDKLGAIVTVNDGKPFDPNPAQCLLESGIKVI 60

Query: 65 CGSHPLELLEDEFCYMIKNPGIYPNNPMVKKALEKQIPVLTEVELAYLVSESLIGITGS 124
  G HPLELLEDE+F N+KNPGIPY+NP++KAL K IPVLTEVELAYL+SE+ +IGITGS
15 Sbjct: 61 TGGHPLELLEDEEFALMVKNPGIYPNSNMIEKALAGIIPVLTEVELAYLISEAPIIGITGS 120

Query: 125 NGKTTITTTMIAEVLNAGGQGLLAGNIGFPASEVVQAANDKTLVMESSPQLMGVKEFR 184
  NGKTTITTTMI EVL A GQ GLL+GNIG+PAS+V Q A DK+TLVMESSPQLMGV+EF
20 Sbjct: 121 NGKTTITTTMIGEVLTAAQGHLLSGNIGYPASQVAQIATDKNTLVMESSPQLMGVQEFH 180

Query: 185 PHIAVITNLMPHLDYHGSFEDYVAAKWNINQMSSDFVLNFMNQGISKEAKTTKATI 244
  P IAVITNLMPH+DYHG FE+YVAAKWNINQ++H++DFVLNFMNQ + K+LA T+At+
25 Sbjct: 181 PEIAVITNLMPHLDYHGLFESYVAAKWNINQMNTAADFVLNFMNQDLVKDLASKTEATV 240

Query: 245 VPFSTTEKVDGAYVQDKQLFYKGENIMSVDDIGVFGSHNVENALATIAVAKLAGISNMVI 304
  VPFST EKVDGAY++D QL+++GE +H+ +IGVFGSHNVENALATIAVAKL G+ NQ I
30 Sbjct: 241 VPFSTLEKVDGAYLEDGQLYFRGEVVMANISIGVFGSHNVENALATIAVAKLRGVNQT 300

Query: 305 RETLSFEGGVKHKRLQSLGKHVHGISFYNDKSTNIIATQKALSGPDNKKVILIAOGLDRGN 364
  +ETLS FGGVHKRLQ + +G+ FYNDKSTNIIATQKALSGPDN+KV+LIAOGLDRGN
35 Sbjct: 301 KETLSAFGGVHKRLQVDDIKGVKFYNDKSTNIIATQKALSGPDNKKVILIAOGLDRGN 360

Query: 365 EFDELIPDITGLKHMVVISGSASVKRAAQKAGVYTSDALDVKRAVHKAYFVAQGGDVL 424
  EFDEL+PDITGLK MV+LG+SA RVKRAA KAGV Y +A D+ DA KAYE+A QGDV+L
40 Sbjct: 361 EFDELIPDITGLKHMVVISGSARVRAADKAGVYVATDIAATRKAYEATQGGDVL 420

Query: 425 LSPANASWDMYKNFEVRGDEFIDTFESLR 453
  LSPANASWDMY NFEVRGD FIDT L+
45 Sbjct: 421 LSPANASWDMYANFEVRGDLFIDTVAELK 449

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 209> which encodes the amino acid sequence <SEQ ID 210>. Analysis of this protein sequence reveals the following:

```

40 Possible site: 25
    >>> Seems to have a cleavable N-term signal seq.

    ----- Final Results -----
45      bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
        bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
        bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

50 RGD motif: 436-438

```

An alignment of the GAS and GBS proteins is shown below:

```

  Identities = 329/451 (72%), Positives = 397/451 (87%)

55 Query: 5 MKTITTFENKKVLVLGLARSGSAAARLLAKLGAIVTVNDGKPFDPNPTAQSLLLEGIKVV 64
  MK I+ P+NKK+L+LGLA+SGRAAA+LL KLG+VTVND KFPD+NP AQ+LLESGIKV+
  Sbjct: 1 MKVISNPFQKKLLILGLAKSGRAAAKLLTKIGALVTVNDGKPFDPNPAQALLLESGIKVI 60

Query: 65 CGSHPLELLEDEFCYMIKNPGIYPNNPMVKKALEKQIPVLTEVELAYLVSESLIGITGS 124
  CGSHP+ELLDE+F YM+KNPGIPY+NPVVK+AL K+IP+LVEVELAY VSE+ +IGITGS
60 Sbjct: 61 CGSHPLELLEDEFCYMVKNPGIYPNNPMVKALAKEIPILTEVELAYVPSEAPIIGITGS 120

Query: 125 NGKTTITTTMIAEVLNAGGQGLLAGNIGFPASEVVQAANDKTLVMESSPQLMGVKEFR 184

```



```

NGKTTTTTMI+VLNAGGQ LL+GNIG+PAS+VVQ A DTLVMESSSFL+GV FR
Sbjct: 121 NGKTTTTTMIADVLNAGGQSALLSGNIGYPASKVQKALAGDTLVMELSSFLGVGNAPR 180

5 Query: 185 PHIAVITNLMPTHLDYHGSFEDYVAAKNIQNMSSDFVLNPNQGISKEIAKIKTATI 244
    PHIAVITNLMPTHLDYHGSFEDYVAAKV IQ QM+ SD+L+LN NQ IS LAKITKAT+
Sbjct: 181 PHIAVITNLMPTHLDYHGSFEDYVAAKNIQNMESDYLLLNANQISATLAKIKTATV 240

Query: 245 VPFSTTEKVDGAYVQDKQLFYKGENIMSVDDIGVPGSHINVENALATIAVAKLAGISNQVI 304
    +PFST + VDGAY+D L++K + I++ D+GVPGSHN+ENALATIAVAKL+GI++ +I
10 Sbjct: 241 IPFSTQKVDGAYLKDGLYFKQAI LAATDLGVPGSHINENALATIAVAKLSGIADII 300

Query: 305 RETLSNFGGVKHLQGLSKVEHGISFYNDKSTNLTATQKALSGFDNTKVIILAGGLDRN 364
    + LS+FGGVKHLQ +G++ I+FYNDKSTNLTATQKALSGFDN+++ILIAGGLDRN
Sbjct: 301 AQLSHFGGVKHLQGVQQLKIDITFYNDKSTNLTATQKALSGFDNRLILIAGGLDRN 360

15 Query: 365 EFDELIPDITGLKHMVVLGESASVKRAAKGAGVTYSALDVRDAVHKAYEVAQQSGDVL 424
    EFD+L+PD+ GLK M++LGESA R+KRAA KA V+Y +A +V +A A++AQ GD IL
Sbjct: 361 EFDLVPDLGLKQMILGESAERKRAANKAEVSYLEARNVAKATELAFKLAQTGDIL 420

20 Query: 425 LSPANASWDMYKNEFVRGDEFIDTFESLRGE 455
    LSPANASWDMY NFEVRGDEF+ TF+ LRG+
Sbjct: 421 LSPANASWDMYNFEVRGDEFIATFDCLRGD 451

```

SEQ ID 208 (GBS305) was expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 51 (lane 11; MW 53.7kDa). It was also expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 56 (lane 3; MW 79kDa).

The GBS305-GST fusion product was purified (Figure 207, lane 8) and used to immunise mice. The resulting antiserum was used for FACS (Figure 270), which confirmed that the protein is immunoreactive on GBS bacteria.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 66

A DNA sequence (GBSx0066) was identified in *S. galactiae* <SEQ ID 211> which encodes the amino acid sequence <SEQ ID 212>. Analysis of this protein sequence reveals the following:

```

35 ROD motif 285-287

Possible site: 60

>>> Seems to have no N-terminal signal sequence
40 INTEGRAL Likelihood = -1.65 Transmembrane 74 - 90 ( 73 - 93)

----- Final Results -----
bacterial membrane --- Certainty=0.1659 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
45 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 213> which encodes the amino acid sequence <SEQ ID 214>. Analysis of this protein sequence reveals the following:

```

50 Possible site: 37

>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood = -1.33 Transmembrane 81 - 97 ( 80 - 100)
INTEGRAL Likelihood = -0.16 Transmembrane 272 - 288 ( 271 - 288)

55 ----- Final Results -----
bacterial membrane --- Certainty=0.1532 (Affirmative) < succ>

```

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

A related sequence was also identified in GAS <SEQ ID 9141> which encodes the amino acid sequence

5 <SEQ ID 9142>. Analysis of this protein sequence reveals the following:

Possible site: 37

>>> Seems to have no N-terminal signal sequence

10 INTEGRAL Likelihood = -1.33 Transmembrane 74 - 90
INTEGRAL Likelihood = -0.16 Transmembrane 265 - 281

----- Final Results -----

15 bacterial membrane --- Certainty=0.1532 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

RGD motif: 286-288

An alignment of the GAS and GBS proteins is shown below:

20 Identities = 249/358 (69%), Positives = 293/358 (81%), Gaps = 1/358 (0%)

Query: 1 MGKKIVFTGGSTVGHVITNLILIPKFIKDGNEVHYIGDKNGIEHEQINQSGLDITFHSIA 60
M KKI+FTGGSTVGHVITNLILIPKFIKDGNEVHYIGDKNGIEH +I +SGLD+TFH+IA
Sbjct: 8 MPKKILFTGGSTVGHVITNLILIPKFIKDGNEVHYIGDKNGIEHEQINQSGLDITFHSIA 67

25 Query: 61 TGKLRVYFSWQNLVDVFKVGVGLQSLAIKLRPQALFSKGGFVSVPFVVAARLLKVPV 120
TGKLRVYFSWQNL+ DVFKV +G+LQS+ I+AKLRPQALFSKGGFVSVPFVVA+LL PV
Sbjct: 68 TGKLRVYFSWQNLADVFKVALGLQLSFLIVAKLRPQALFSKGGFVSVPFVVAARLLKVPV 127

30 Query: 121 FVHESDLNSGLANKIAYKFATIMYTFPQSKOLIKTHIGAVTKVM-DCKSEFNIDTIS 179
F+HESD SMLANKIAYKFAT MYTFPQ L K KH+GAVTKV D + E+T L +
Sbjct: 128 FVHESDRSGLANKIAYKFATIMYTFPQKQDLKSKVHIGAVTKVFKDANQMPSTQEA 187

35 Query: 180 IREAFDNLKTLFLFIGGSAGAKVFNDFITQTPLEEKYNVINISGDSINRLKNLRYVD 239
+KE F +LKTLLFIGGSAGA VFN FI+ PEL+++YN+INI+GD LN L +LYRVD
Sbjct: 188 VREYFSDRLKTLFLFIGGSAGAHVFNQFISDHPFLKQRYNIINITGDPHLNLSHLYRVD 247

40 Query: 240 YVTDLYQPLMADLVVVTROGSGNTIFELVAMKHLHLIPLGSEASRGDQLENAAYPEEK 299
YVTDLYQPLM +AD+VVTROGSGNT+PEL+AM KLHLI+PLG+EASRGDQLENA YFE+G
Sbjct: 248 YVTDLYQPLMADLVVVTROGSGNTIFELVAMKHLHLIPLGSEASRGDQLENAAYPEEK 307

45 Query: 300 YALQLPESELNINTLEKQINLLISNSESYEKNMSOSSEIKSQDEFFYLLIDDMKVTK 357
YA QL E L + + + + L + YE M + EI+ S D FY LL D+ + K
Sbjct: 308 YALQLQEPDLTLNFPQAMADLFEHQADYEATLATKEIQSPDFYLLRADISSAIK 365

SEQ ID 212 (GBS306) was expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 51 (lane 12; MW 43kDa). It was also expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 56 (lane 4; MW 68kDa).

GBS306-GST was purified as shown in Figure 207, lane 9.

50 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 67

A DNA sequence (GBSx0067) was identified in *S. agalactiae* <SEQ ID 215> which encodes the amino acid sequence <SEQ ID 216>. This protein is predicted to be cell division protein DivIB. Analysis of this protein
55 sequence reveals the following:

Possible site: 58

-126-

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -14.33 Transmembrane 103 - 119 (96 - 124)

5 ----- Final Results -----
 bacterial membrane --- Certainty=0.6731 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

10 The protein has homology with the following sequences in the GENPEPT database:
 >GP: AAC95451 GB: AF068902 cell division protein DivIB [Streptococcus pneumoniae]
 Identities = 119/396 (30%), Positives = 214/396 (53%), Gaps = 38/396 (9%)

15 Query: 3 KKS DTPKEEVV- L TEWQKRNLEFLKRRKDEE --- RQKRNELRLDKRS ---- KLN 53
 KK D EE+ L+EWQKN E+LKK+ E+E K+ +R+ S K+
 Sbjct: 5 KKNEDKEILLEELSELSEWQKRNQYELKKKKEEAALEKEKEKQARMGSEBSEKQD 64

Query: 54 ISSPEEPQNTTKIKLHFPKIS ----- RPKIEKKQKKEKIVNSLAKTNR --- 97
 S + +++ K+ K++ P+ ++K+++K ++ A +
 20 Sbjct: 65 QBSRTDQDSBSAKRESEKVSASADKEKEKEEPEKKEEQQDKLKKKATKPKAKA 124

Query: 98 ----- IRTAPIFVVAFLVILVSFLLTPFSKQKTTITVSGNQHTPDILIEKTIQKND 150
 +R I + L+++VS +LL+P++ K I V G T D + + + IQ +D
 Sbjct: 125 KIPGIHLFAFTILFPLSLLILVSAYLSPYATMKDIRVEGTQVTALDIRQASIGQSD 184

25 Query: 151 YPFSLIFPKHKAIEQRILAAEDVWVKTAQMTYQFPNKFHQVQENKIIAYARTKQGVQPVLE 210
 Y +L+ E+++ + + WV++AQ+ YQFP KF I+V+E I+AY + + + P+L
 Sbjct: 185 YTNILLDLKKYKQIKS-NYVWESAGLVQYPTFKTIFKVEKYDIVAYYISGENYPIIS 243

30 Query: 211 TGG-KADPVNSSELKPKHFLTINLDKEDSKILLIKDLKALDPDLISIQVLSIADSKTTPD 269
 +G+ + V+ + LP+ +L++ + + IK+ +L + P+L + IQ + LA SK T D
 Sbjct: 244 SQGLSETSSVLSNLSLPETYLVSFLNDSQIKVPVSELAQISPELKAATQKVLAKSPILS 303

35 Query: 270 LLLLDMDHGNISIRIPLSKFKERLPFYKQIKKIKKEPSIVDMVGVTYTTINIESTPVKAE 329
 L+ L M+D + + +PLS+ ++LP+Y +IK L EPS+VIME G+Y+ T + E
 Sbjct: 304 LIRLTWNDSQVILVPLSEMSKKLFPYYSKIPQLSEPSVDMIEAGTYSYTVADKLINVEE 363

Query: 330 DTKNKSTDKTQTQNGQVAENSQGGTNNSTNCCGQQ 365
 K ++ + Q E + Q SN NQ Q+
 40 Sbjct: 364 KAKQEAKEAEKQGE---EEQKKQEEESNRNQITQR 395

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 217> which encodes the amino acid sequence <SEQ ID 218>. Analysis of this protein sequence reveals the following:

Possible site: 59
 45 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -9.45 Transmembrane 106 - 122 (102 - 125)

----- Final Results -----
 bacterial membrane --- Certainty=0.4779 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

50

An alignment of the GAS and GBS proteins is shown below:

Identities = 152/381 (39%), Positives = 232/381 (59%), Gaps = 14/381 (3%)

55 Query: 4 KKS DTPKEEVVILTEWQKRNLEFLKRRKDEE RQKRNELRLDKRS KLN ISSPEEP --- 60
 K + +++VLTEWQKRN+EFLKK+K+ EE+K++ EKL DK+++ + E
 Sbjct: 3 KDKSKQSDDKLVLTENQKRNIEFLKRRKQAEERKKLEKLLSDKKAQQQANASRAVEL 62

60 Query: 61 --QNTTKIKKLHFPKISRPKIEKK--QKKEKIVNSLAKTNRI RTAPIFVVAFLVILVSF 116
 T +++ S+PK KK Q KEK +A ++ P+ + A L++ VS+P
 Sbjct: 63 KTDKTDQSEIESSTTSKPKKTKVKVQPKESATQIAFQ---KSLPVLGALLMAYSTP 119

Query: 117 LLTPFSKQKTTITVSGNQHTPDILIEKTIQKNDYFSSLIFPKHKAIEQRILAAEDVWVKTA 176

```

      ++TP+SK+K +V GN T D LI+ ++ +DY+ +L+ E+ + WVK+
Sbjct: 120 MITPYSKKEFSVRGNHQTNLDELKASKVKASDYMLTLTSPGQVERPIILRTIPWKEV 179

5 Query: 177 QMTYQFPNKHFLQVQENKIIAYAHFKQYQPVLETGKKADPVNSBLPKHFLATINLQDEK 236
      ++YQFPN P V E +IIAYA + G+Q+LE GK+ D V +SRLEPK FL +NL E
Sbjct: 180 HLSYQFPNKHFLNVIEFELIAYAQVENGFPQELIENGRKVDKVRASLEPKSFLIATLQDEK 239

Query: 237 SIKLLIKDLKALDPLDISIQLVISLADSKITPDLLILLMDHGNSTRIPLSKFKERLEPFYK 296
      +I+ L+K L L L+ I+ +SLA+SKIT DLLL+MHGDN +R+P S+ +LP+Y+
10 Sbjct: 240 AIQQLVQLITLTPKLVKNIKSVSLANSKITADLLILRMHDCNVVRVPSQLATLQPYQ 299

Query: 297 QIKKNLKEPSIVDMVEGVYTTTINTIESTPVKAEDTNKSKTDCTQNGQVAKSSQQQTMN 356
      ++KKNL+ SIVDMRVG+YTTT IE+ P + +DK +G+ Q QT+N
15 Sbjct: 300 KLEKNLENDIVDMVEGVYTTTQIEENQPEVPLTPBQNAADKEGDKRGE---HQEQTMN 355

Query: 357 SNTNQQGQQIATBQAPNIQNV 377
      + O + P+P+ V
Sbjct: 356 DSETFANQSSPQQTPPSPETV 376

```

- 20 SEQ ID 216 (GBS85) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 17 (lane 10; MW 45.2kDa).

The GBS85-His fusion product was purified (Figure 105A; see also Figure 193, lane 5) and used to immunise mice (lane 1 product; 20µg/mouse). The resulting antiserum was used for Western blot (Figure 105B), FACS (Figure 105C), and in the *in vivo* passive protection assay (Table III). These tests confirm

25 that the protein is immunoaccessible on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 68

A DNA sequence (GBSx0068) was identified in *S.galactiae* <SEQ ID 219> which encodes the amino acid

30 sequence <SEQ ID 220>. This protein is predicted to be cell division protein FtsA (ftsA). Analysis of this protein sequence reveals the following:

```

Possible site: 56

>>> Seems to have an uncleavable N-term signal seq
INTEGRAL Likelihood = -3.19 Transmembrane 322 - 338 ( 321 - 338)

35 ----- Final Results -----
      bacterial membrane --- Certainty=0.2275 (Affirmative) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
40      bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AC95439 GB:AF068901 cell division protein FtsA [Streptococcus pneumoniae]
Identities = 292/457 (63%), Positives = 366/457 (79%), Gaps = 1/457 (0%)

45 Query: 1 MARNQFTGLDITGTSIKVLVAEFIANENNVIGSVNVPSSGVKDGIIIDIRAAATAIKSA 60
      MAR GFPTGLDITGTS+KVLVAE E+NVIGSVN S GVKDGI+DI+MAATAIK A
Sbjct: 1 MAREGFPTGLDITGTSVKVLVAQRNGELNVIGSVNAKSGVKDGIIVDIDAAATAIKSA 60

50 Query: 61 VQAEKAGITIDKINGVLENNLQIEPTQGMIPVNPESKEIKDEBDSVSVKALTSKST 120
      + QAEEKAGI+I +NVGLP NLLQ+EPTQGMIPV +++KEI D+DVE+VVKALTSK+T
Sbjct: 61 ISQAEKAGISIKSVNVGLFGNLLQVEPTQGMIPVTSUTKEITDQVENNVKALTSKST 120

Query: 121 PEREVISLILFLEPIDVGGQGIROPGRMGVLEMRGLIYTGPTTLINLRKTVRAGIKV 180
      P+REVI+ IP EFIDVGGQGIROPGRMG+LEMRGL+YTGPT TLINLRKTVRAG+V
55 Sbjct: 121 POREVITTFEPEPIDVGGQGIROPGRMGVLEMRGLIYTGPTTLINLRKTVRAGVQV 180

```

-128-

Query: 181 EHVVIAPLALAKSVLNEGGEREPGATVIDMGGQTTVASMRNQLQYTNIIYSEGGSDVVTKD 240
 E+V+I+PLA+ +SVLNEGGEREPGATVIDMG GQTTVA++RNOELQ+T+I EG DVVTKD
 Sbjct: 181 ENVIISPLAVQSVLNEGGEREPGATVIDMGAQTTVATIRNQLQFTHILQEGGSDVVTKD 240

Query: 241 ISKVLRTTVELAEALKPNFQGANVEASTSDTVQNVVONEPVEITESYLSQIISGRIR 300
 ISKVL+T+ ++AR LK N+G+A AS +T QV V+G E VE+TE+YLS+IIS RI+
 Sbjct: 241 ISKVLKTSRKLAEGLKLNFGYAPPLAS-KETQVEVIGVEAVEVTEAYLSEIISARIK 299

Query: 301 QILRHVQKDLGRRLDLPGGIILVGGGAIMPGVVEVAQQIFQTRVKLHVPAQVGIRNMP 360
 ILE +KQ+L R RLDDLPGGI+L+GG AI+PG+VE+AQ++PG RVKL+VPAQVGIRNP
 Sbjct: 300 HILEQIKQELDRRLDLPGGIVLIGGNAILFGMVLEAEVFGVRVKLYVPAQVGIRNPA 359

Query: 361 FANVISVDVYGMMSVDDILAQHAVTGERMLRHKFPDVPDYKEKINMTMTPSYEPLTSS 420
 FA+VIS+ ++ G ++EV+++AQ A+ G+ L H+P+ F + +
 Sbjct: 360 FAHVISLSFAGQLTEVNLLAQAIGKENDLSHQPISEFGMLQKTAQFVQSTPVQAPAPAF 419

Query: 421 EDSNLEPIRARENAQETPEPKANIGERIRGIFGSMFD 457
 E + P + Q+ ++ K + +R RG+ GSMFD
 Sbjct: 420 ESEVAPTEINWADPQASCNKPKLADRFRLIGSMFD 456

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 221> which encodes the amino acid sequence <SEQ ID 222>. Analysis of this protein sequence reveals the following:

Possible site: 55

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -3.35 Transmembrane 313 - 329 (312 - 329)

----- Final Results -----

bacterial membrane --- Certainty=0.2338 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP:AA095439 GB:AF068901 cell division protein FtsA [Streptococcus pneumoniae]
 Identities = 299/448 (66%), Positives = 368/448 (81%), Gaps = 4/448 (0%)

Query: 1 LDIGTSSIKVLVAEFISGEMNVIGSVNPSTGVKDGIIIDIEAATAIKTAVQAEIRAG 60
 LDIGTSS+KVLVAE +GE+NVIGVSN S GVKDGII+DI+AAATAIK+A+ QAEIRAG
 Sbjct: 10 LDIGTSSIKVLVAEQRXGELNVIGSVNNAKSGVKDGIIVDIDRAATKASASQAEIRAG 69

Query: 61 MTIEKVNVLGPANLLQIEPTQGMIPVPSSEKRIKDEVDVSVKSALTSTKSTPEREIVSLV 120
 ++I+ VNVGLP NLQ+EQPTQGMIPV S++KEI D+DV++VVSALTSTK+TP+REVI+ +
 Sbjct: 70 ISIKSVNVLGPNLLQVSEPTQGMIPVTSYDKRIKDEVDVSVKSALTSTKSTPEREIVTFL 129

Query: 121 PEEFIVDGFQGIIRDPRGMGIRLDMRGLIYTGPTILHNLKRTVERAGIKVNIISPLA 180
 PEEFIVDGFQGIIRDPRGMG+RLDMRGL+YTPG TILHNLKRTVERAG++VEN+IISPLA
 Sbjct: 130 PEEFIVDGFQGIIRDPRGMGVRLEMRGLIYTGPTILHNLKRTVERAGQVENVISPLA 189

Query: 181 MAKTIILNEGGEREPGATVIDMGGQTTVASMRAQLQYTNIIYARGEGSYITKDISKVLKTS 240
 M++++LNEGGEREPGATVIDMG GQTTVA++R QELQ+T+I EGG+Y+TKDISKVLKTS
 Sbjct: 190 NVQSVLNEGGEREPGATVIDMGAQTTVATIRNQLQFTHILQEGGSDVYTRDISKVLKTSR 249

Query: 241 ALAEALKFNFOQAELSKASITETVKVDVVGSRPVSVTERYLSEIISARIRHILDRVKQD 300
 +AR LK N+G+A AS ET +V+V+G E VEUTE YLSEIISARI+HIL++RQ+
 Sbjct: 250 LABGLGKLNFGYAPPLAS-KETQVEVIGVEAVEVTEAYLSEIISARKHILEQIKR 308

Query: 301 LERGRLLDLPGGIVLIGGGAIMPVVEIAQELFGVTKLHVPAQVGIRNMPFNVISLVE 360
 L+R RLDDLPGGIVLIGG AI+PG+VE+AQ+PGV VKL+VPAQVGIRNP F++VISL E
 Sbjct: 309 LDRRLDLDLPGGIVLIGGNAILFGMVLEAEVFGVRVKLYVPAQVGIRNPAFAHVISLS 368

Query: 361 YVGMMSVDVLAQTAVSGELLRRKPIDFSQGESYLPDYDDSRPESTIGYEQ-- --ASQ 417
 + G ++EV++LQ A+ GE L +PI F G + S + E + ++
 Sbjct: 369 FAGQLTEVNLLAQAIGKENDLSHQPISEFGMLQKTAQFVQSTPVQAPAPAVEVEFAPTE 428

Query: 418 TAYDSQVPSDPKQKISERVRGIFGSMFD 445
 D Q S K K++R RG+ GSMFD
 Sbjct: 429 PMADPQASQNKPKLADFRGLIGSMFD 456

5 An alignment of the GAS and GBS proteins is shown below:

Identities = 349/456 (76%), Positives = 402/456 (87%), Gaps = 19/456 (4%)

Query: 10 LDIGTSSIKVLVAEFTANEMVIGSVPSGVKDGIIIDIAAATAIKVAKQAEKAG 69
 LDIGTSSIKVLVAEFT+ EMEVIGSVPS+GVKDGIIIDIAAATAIK AV+QAEKAG
 10 Sbjct: 1 LDIGTSSIKVLVAEFTISGRMEVIGSVPSGVKDGIIIDIAAATAIKVAKQAEKAG 60

Query: 70 ITIDKINVLGPNLQLIEPTQGMIPVNSEKKEIKDEDVSVVKSALTKSITPEREVLISL 129
 +TI+K+NVGLPNLQLIEPTQGMIPV+SEKKEIKDEDV+SVVKSALTKSITPEREVLISL+
 15 Sbjct: 61 MTIEKINVLGPNLQLIEPTQGMIPVPSSEKKEIKDEDVSVVKSALTKSITPEREVLISL 120

Query: 130 PLEFIVDGFQGIIRPRGMGIRLEMRLIYTGPTILHNLRTKTVRAGIKVEHVIAPLA 189
 P E FIVDGFQGIIRPRGMGIRLEMRLIYTGPTILHNLRTKTVRAGIKVE++I+PLA
 15 Sbjct: 121 PEEFIVDGFQGIIRPRGMGIRLEMRLIYTGPTILHNLRTKTVRAGIKVEHIIISPLA 180

Query: 190 LAKSVLNBGEREPGATVIDMGGGQTTVASMRLQYINIIYSESGDYVTKDISKVLRTTV 249
 +AK++LNBGEREPGATVIDMGGGQTTVASMRLQELQYINIIY+EG +Y+TKDISKVL+T++
 20 Sbjct: 181 MAKTYLNBGEREPGATVIDMGGGQTTVASMRLQELQYINIIYABGGHYTKDISKVLRTTV 240

Query: 250 EIAEALKFNFGQAQVKEASTSDIVQVNVGNPEPVKITEGYLSQIISGRIRQLHVKQD 309
 IAEALKFNFGQA +EAS ++TV+V+VVG+BEPEV+TE YLS+IIS RIR IL+ VKQD
 25 Sbjct: 241 AIAEALKFNFGQASISEASTYETVKVNVVGSSEPVKTEGYLSIISARILHILRVKQD 300

Query: 310 LGRGRLLDLPGGIILVGGGAIMPGVVEVAQQIFGTRVKLHVFNGWIRNPFNFANVISVD 369
 L RGRLLDLPGGI+L+GGGAIMPGVVE+AQ+IFG VKLHVFNGWIRNPFNFANVIS+V+
 30 Sbjct: 301 LERGRLLDLPGGIILVGGGAIMPGVVEIAQIFGTVTKLHVFNGWIRNPFNFANVISLVE 360

Query: 370 YVGMSEVDIIAQAHTVDENLRHKFVDF-----DYKEKNTNTMTMPSEYPTSSNE 421
 YVGMSEVD++AQ AV+G+E+LR KF+DF DY + ST+ Y + + +
 35 Sbjct: 361 YVGMSEVDVLAQTAVSGRELLRRKPIDFSGQESYLPDYDERRPESTIGYEQASQTAY 420

Query: 422 DSNLEPIRARENAQEPTPEKANIGERIRGIFGSMFD 457
 DS Q P+PK I ER+RGIFGSMFD
 Sbjct: 421 DS-----QVPSDPKQKISERVRGIFGSMFD 445

40 SEQ ID 220 (GBS73) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 17 (lane 5; MW 47.8kDa). It was also expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 20 (lane 5; MW 70.1kDa).

GBS73-GST was purified as shown in Figure 197, lane 7.

The GBS73-His fusion product was purified (Figure 103A) and used to immunise mice (lane 1 product; 20µg/mouse). The resulting antiserum was used for Western blot (Figure 103B), FACS (Figure 103C) and in the *in vivo* passive protection assay (Table III). These tests confirm that the protein is immunoaccessible on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

50 **Example 69**

A DNA sequence (GBSx0069) was identified in *S.galactiae* <SEQ ID 223> which encodes the amino acid sequence <SEQ ID 224>. This protein is predicted to be cell division protein FtsZ (ftsZ). Analysis of this protein sequence reveals the following:

Possible site: 56

-130-

>>> Seems to have a cleavable N-term signal seq.

INTEGRAL Likelihood = -1.97 Transmembrane 117 - 133 (117 - 133)

5 ----- Final Results -----

bacterial membrane --- Certainty=0.1786 (Affirmative) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

10 The protein has homology with the following sequences in the GENPEPT database:

>GP:AAC95440 GB:AF068901 cell division protein FtsZ [Streptococcus pneumoniae]

Identities = 327/426 (76%), Positives = 363/426 (84%), Gaps = 7/426 (1%)

15 Query: 1 MVFSFDTASVQGAIVKIVGVGGGGGNAINRMIDEGVAGVEFIAANTDIALSSSKAETVI 60

M FSTDTA+ QGAVIKIVGVGGGGGNAINRM+DEGV GVEFIAANTD+QALSS+KAETVI

Sbjct: 1 MTFSFDTAAAGAVIKIVGVGGGGGNAINRMVDGVEFIAANTDQALSSSTKAETVI 60

Query: 61 QLGPXILTRGLGAGGGQPEVGRKAAEESERVLTALTGADMFVITAGMGGGSGTGAAPVIAR 120

QLGPXILTRGLGAGGGQPEVGRKAAEESER LTEA++GADMFVITAGMGGGSGTGAAPVIAR

20 Sbjct: 61 QLGPXILTRGLGAGGGQPEVGRKAAEESERVLTALTGADMFVITAGMGGGSGTGAAPVIAR 120

Query: 121 IAKSLGALTVAVITRPFQPEGMKRSNFAIEGIELREBQVDTLLIISNNLLEIVDKTKPL 180

IAK LGALT V+TRPFQPEG+KR FA+EG I+LRE VDTLLIISNNLLEIVDKTKPL

25 Sbjct: 121 IAKDLGALTVGIVTRPFQPEGSKRGQFAVEGINQLREHVDTLIISNNLLEIVDKTKPL 180

Query: 181 LEALSEADNVLROGVGGITDLITNPGLINLDFADVTKVMANKGNALMGIGSGSERITE 240

LEALSEADNVLROGVGGITDLITNPGLINLDFADVTKVMANKGNALMGIGSGSERE+ E

30 Sbjct: 181 LEALSEADNVLROGVGGITDLITNPGLINLDFADVTKVMANKGNALMGIGSGSERIVE 240

Query: 241 AARKAIYSPLLETTIDGSEADVINVTGGMDMTLIEAREASEIVSQAGKGVNIWLGTSID 300

AARKAIYSPLLETTIDGSEADVINVTGG+D+TL IEAREAS+IV+QAG+GVNIWLGTSID

35 Sbjct: 241 AARKAIYSPLLETTIDGSEADVINVTGGDLTLIEAREASQIVNQAGQGVNIWLGTSID 300

Query: 301 MDMKDEIRIVTVATGVRIDKTNQSGFTTSAPTNQAFSEQRSTSNENFDRRGNFDMTESR 360

M+DEIRIVTVATGVR+D+ +V +TN + + + S+ PDR +PDM E+

40 Sbjct: 301 ESMRDEIRIVTVATGVRQDRVEKVFQARSATNYRETVPKPAHSH-GFDR--HFDMAETA 357

Query: 361 EMPTQQNQPHAQNCQSSAPGNMDLRDNI SRPTGELDSKLSMETTSSENDMDDELTTP 420

E+P Q P Q+SAPG+MDLRR++I R T+ + D +DEL+TP

45 Sbjct: 358 ELPKQ--NPRRLPTQASAPGMDLRRESIVRTDSVVSFVERFEAPISQD--EDELTP 413

Query: 421 PFFKNR 426

PFFKNR

Sbjct: 414 PFFKNR 419

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 225> which encodes the amino acid sequence <SEQ ID 226>. Analysis of this protein sequence reveals the following:

Possible site: 56

>>> Seems to have a cleavable N-term signal seq.

INTEGRAL Likelihood = -1.81 Transmembrane 117 - 133 (117 - 133)

----- Final Results -----

bacterial membrane --- Certainty=0.1723 (Affirmative) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 372/439 (84%), Positives = 391/439 (88%), Gaps = 13/439 (2%)

60 Query: 1 MVFSFDTASVQGAIVKIVGVGGGGGNAINRMIDEGVAGVEFIAANTDIALSSSKAETVI 60

M FSFDTAS+GGA+IVKIVGVGGGGGNAINRMIDEGVAGVEFIAANTDIALSSSKAETVI

Sbjct: 1 MAFSFDTASIGGAIVKIVGVGGGGGNAINRMIDEGVAGVEFIAANTDIALSSSKAETVI 60

Query: 61 QLGPKLTRGLGAGGQPEVGRKKAARESESVLITRALTGDMVFTITAGMGGSGTGAAAPVIAR 120
 Sbjct: 61 QLGPKLTRGLGAGGQPEVGRKKAARESEELITRALTGDMVFTITAGMGGSGTGAAAPVIAR 120

5 Query: 121 IAKSLGALTAVVITRPFGEFGNKRNFALBGIQELREQVDTLITISNNNLEIVDKKTFPL 180
 IAKSLGALTAVV+TRPFGEFGNKR NFALBGI+ELREQVDTLITISNNNLEIVDKKTFPL 180
 Sbjct: 121 IAKSLGALTAVVITRPFGEFGNKRNFALBGIQELREQVDTLITISNNNLEIVDKKTFPL 180

10 Query: 181 LEALSEADNVLRQGVGIDTLITNFGLINLDPADVKTVMANKGNALMGIGSGSERITE 240
 LEALSEADNVLRQGVGIDTLIT+FGLINLDPADVKTVMANKGNALMGIGSGSERITE 240
 Sbjct: 181 LEALSEADNVLRQGVGIDTLITNFGLINLDPADVKTVMANKGNALMGIGSGSERITE 240

15 Query: 241 AARKAIYSPILLETTIDGAEDVIVNVITGMDMTLTAEEASEIVSQAGKGVNIWLGTSID 300
 AARKAIYSPILLETTIDGA+DVIVNVITG+DMTLEAEASEIV QAAG+GVNIWLGTSID 300
 Sbjct: 241 AARKAIYSPILLETTIDGAEDVIVNVITGMDMTLTAEEASEIVSQAGKGVNIWLGTSID 300

20 Query: 301 MDKDEIRVITVATGVRKDKTNQVSGF---TTSAPTN-----QAPSERQSTENSNFD 349
 MKD+IRVITVATGVR++K QVSGF T TN A + + + FD 349
 Sbjct: 301 DTMKDDIRVITVATGVRQEKAEQVSGFRQPTFTQTNAQCVAGAQVASDQAKVSGQPF 360

25 Query: 350 REGN--FDMTESREMPQQNQPHANQQQSFAFGNWLRRDNISRPTBGEGLKLMSTTF 407
 RR N FDM ESRE+P+ Q NQ Q SAFGNWLRRDNISRPTBGEGL+ L+MSTTF 407
 Sbjct: 361 RRSNFDPMGSRERIPSAKQVLSNNHNNQGSFAFGNWLRRDNISRPTBGEGLNHLNMSTF 420

30 Query: 408 SBNDDMDDELSTPPFFPKNR 426
 S NDD DDELSTPPFFPKNR 426
 Sbjct: 421 SANDDSDDELSTPPFFPKNR 439

SEQ ID 224 (GBS163) was expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 28 (lane 7; MW 44kDa). It was also expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 34 (lane 4; MW 69kDa).

The GBS163-GST fusion product was purified (Figure 114A; see also Figure 198, lane 11) and used to immunise mice (lane 1 product; 20µg/mouse). The resulting antiserum was used for Western blot (Figure 114B), FACS and in the *in vivo* passive protection assay (Table III). These tests confirm that the protein is immunoreactive on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 70

A DNA sequence (GBSx0070) was identified in *S. galactiae* <SEQ ID 227> which encodes the amino acid sequence <SEQ ID 228>. Analysis of this protein sequence reveals the following:

Possible site: 21

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2750 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AACS5442 GB:AF068901 Ylm8 [Streptococcus pneumoniae]
 Identities = 140/223 (62%), Positives = 177/223 (78%)

Query: 2 MNLEQNKTAIFDNVSKLALAGRAHESVHIVAVTKYVNCQTEALLIRFGVNHIGENRVDK 61
 MN++EN +F V++ +L A R SV +AVTKYV+ T EAL+ GV+HIGENRVDK 61
 Sbjct: 1 MNVKTENELVFREVAEASLSAHRREGSGSVSIAN+KYVDVPTREALLPLGVHIGENRVDK 60

-132-

Query: 62 FLEKYQALKDEKLTWHLIGSLQRRKVKDVINYVDYFHALDSVKLA AEIQHAKLIKCF 121
 FLEKY+ALKD +TWHLIG+LQRRKVKDI YVDYFHALDSVKLA EIQK + ++IKCF
 Sbjct: 61 FLEKYALKDRDVTWHLIGTLQRRKVKDVIOYVDYFHALDSVKLA EIQRSDRVKCF 120

Query: 122 QVNISREDSKHGFTIEQIDDAI NLSRYDKIELIGINTMAPLKATKEIISIFESTESLR 181
 QVNIS+E+SKHGF+ E++ + L ++R DKIE +G+MTMAP +A+ E++ IF+ + L+
 Sbjct: 121 QVNISKEESKHGFSRELELEILPELARLDKIEYVGLINTMAPFEASSEOLKEIFKAQDLQ 180

Query: 182 KRLQARNIERMPFTELSGMSRDYDIAIQNGSTFVRIGTSFFK 224
 + +Q + I MP TELSGMSRDY AIQ GSTFVRIGTSFFK
 Sbjct: 181 REIQEKQIPMMFTELSGMSRDYKRAIQGSTFVRIGTSFFK 223

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 229> which encodes the amino acid sequence <SEQ ID 230>. Analysis of this protein sequence reveals the following:

Possible site: 20

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm	---	Certainty=0.2451(Affirmative)	< succ>
bacterial membrane	---	Certainty=0.0000(Not Clear)	< succ>
bacterial outside	---	Certainty=0.0000(Not Clear)	< succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 133/222 (59%), Positives = 164/222 (72%)

Query: 2 MNLQENKTAIFDNVSKLALQGRAHESVHIVATKYVNCQTTTEALIRTGVNHIENRVDK 61
 M+L NK IP+ + A R ++SV ++AVTKYV+ LI G+ HI ENRVDK
 Sbjct: 1 MDLLTNKKKIFETIRLSTEANNRTNDSVSVIAVTKYVDSTIAGQLIEAGIEHIAENRVDK 60

Query: 62 FLEKYQALKDEKLTWHLIGSLQRRKVKDVINYVDYFHALDSVKLA AEIQHAKLIKCF 121
 FLEKY ALK + WHLIG+LQRRKVK+VINYVDYFHALDSV+LA EI K A +KCF
 Sbjct: 61 FLEKYDALKMPVKHHLIGTLQRRKVKDVINYVDYFHALDSVRLALEINKRADHPVKCF 120

Query: 122 QVNISREDSKHGFTIEQIDDAI NLSRYDKIELIGINTMAPLKATKEIISIFESTESLR 181
 QVNIS+E+SKHGF I +ID+A+ I + +KI+L+G+MTMAP A+KE I +IF + LR
 Sbjct: 121 QVNISKEESKHGFMISSEIDEALGEIGDMEKIQLVGLINTMAPANAKESITITFRQANQLR 180

Query: 182 KRLQARNIERMPFTELSGMSRDYDIAIQNGSTFVRIGTSFF 223
 K LO + + MPTELSGMS DY IAIQ GSTF+RIG +FF
 Sbjct: 181 KNLQLKKRIKMPFTELSGMSRDYPIAIQNGSTFIRIGRAFF 222

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 71

A DNA sequence (GBSx0071) was identified in *S.agalactiae* <SEQ ID 231> which encodes the amino acid sequence <SEQ ID 232>. This protein is predicted to be YlmF. Analysis of this protein sequence reveals the following:

Possible site: 58

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm	---	Certainty=0.2194(Affirmative)	< succ>
bacterial membrane	---	Certainty=0.0000(Not Clear)	< succ>
bacterial outside	---	Certainty=0.0000(Not Clear)	< succ>

A related GBS nucleic acid sequence <SEQ ID 9617> which encodes amino acid sequence <SEQ ID 9618> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```

5 >GP:AAC95442 GB:AF068901 YlmF [Streptococcus pneumoniae]
  Identities = 86/200 (43%), Positives = 120/200 (60%), Gaps = 25/200 (12%)

Query: 5 MALIKDRFDKIISYFDITDVSENEVHVQRRTSVQRDSRAATAQASQSRHMTNSAREMI 64
M+LKDRFD+ I YP T+D + +R +RD T+ +SQ + + +
10 Sbjct: 1 MSLKDRFRFDIDYF-TRDEDSLSLYE-----KRDEPVFTSVNSSQRPALFHNQFSQA 52

Query: 65 GSRPRITYTDENRQERQVRQVDNAYQATPRVONKDSVRQQRBOVTIALKYPKRYBDAQE 124
G++ T RQ+ + N Q+AT ++V I ++YPRKYBDA E
Sbjct: 53 GIKENNITRLHARQQ-----ELANQSRAT-----DKVIIDVRYPRKYBDATE 95

15 Query: 125 IVDLLIVNECVLIDFQYMLDAQARRCLDYIDGASRVLYGSLQKVGSMFLLTPTANVVDI 184
IVDLL NE +LIDFQYM + QARRCLDY+DGA VL G+L+KV S+M+LITP NV+V++
Sbjct: 96 IVDLLAGNESILIDFQYMTVEVQARRCLDYIDGACHVLAGNLKVASTWYLLTPTVNVIVNV 155

Query: 185 ESMNIPKIQGTSFDFDMKR 204
E++ +P Q+ F FDMKR
20 Sbjct: 156 EDIRLPDEDQQGSGFDFDMKR 175

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 233> which encodes the amino acid sequence <SEQ ID 234>. Analysis of this protein sequence reveals the following:

```

25 Possible site: 49
>>> Seems to have no N-terminal signal sequence
    INTEGRAL Likelihood = -0.64 Transmembrane 142 - 158 ( 142 - 158)

----- Final Results -----
30 bacterial membrane --- Certainty=0.1256(Affirmative) < succ>
    bacterial outside --- Certainty=0.0000(Not Clear) < succ>
    bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the databases:

```

35 >GP:AAC95442 GB:AF068901 YlmF [Streptococcus pneumoniae]
  Identities = 82/219 (37%), Positives = 113/219 (51%), Gaps = 46/219 (21%)

Query: 5 MAPKDTFNKIMISYFDITDEVNEVEDVAASTDNVIP--RSQQSVRASSHPKQSPRNHHVQQ 62
M+ KD F++ I YF DE D+P + + V S + QSP Q
40 Sbjct: 1 MSLKDRFRFDIDYFTEDE-----DSSLPYEKRDPEVFTSVNSSQRPALFHNQF 48

Query: 63 DHQARQEQVTSQMEHPKHGTSERYQQSQPKGEHVMVDRKRMSTSSIANRRBQVQQSTC 122
A ++E ++H + +AN Q
Sbjct: 49 SQSAGTKENNITRLHARQ-----QELAN-----QSQRA 76

45 Query: 123 SDQTTIALKYPKRYEDAQEIVDLLIVNECVLIDFQYMLDAQARRCLDYIDGASVLYGSL 182
+D+ I ++YPRKYEDA EIVDLL NE +LIDFQ+M + QARRCLD+DGA VL G+L
Sbjct: 77 TDKVIIDVRYPRKYBDATEIVDLLAGNESILIDFQYMTVEVQARRCLDYIDGACHVLAGNL 136

50 Query: 183 QKVGSMMYLLAPNSVNVIEEMTPIHTTDIGDFDFDMKR 221
+KV S+MYLL P NV VN+E++ +P Q F FDMKR
Sbjct: 137 KKVASTWYLLTPTVNVIVNVEDIRLPDEDQQGSGFDFDMKR 175

```

An alignment of the GAS and GBS proteins is shown below:

```

55 Identities = 118/222 (53%), Positives = 145/222 (65%), Gaps = 17/222 (7%)

Query: 1 MEGNMAIKDRFDKIISYFDITDVSENEVHVQRRTSV---QRDSRAATAQAS----- 50
ME MA KD F+K+ISYFDITD+V+E E +V Q+ RA++ +
60 Sbjct: 1 MENTKAFKDTFNKIMISYFDITDEVNEVEDVAASTDNVIPRSQQSVRASSHPKQSPRNHHV 60

Query: 51 QRSHMTNSAREEMIGSRPRITYTDPNKRQRQRVQR---DNAYQATPRVONKDSVRQQR 106

```

```

      Q+ H   S E+      P+ T+  Q+ Q +      D  + +T + N+  QQ
Sbjct: 61 QQDHQARSRQSPQSRQMHPKHGTSERYYQSQSPKEGHMVDRKKRMSTSSIANRRREQYQQS 120

5  Query: 107 ---EQVTIALKCYPRKYEDAQEIVDLLIVNECVLIDFQYMLDAQARRCLDIYIDGASRVLYG 163
      +Q TIALKCYPRKYEDAQEIVDLLIVNECVLIDFQ+MLDAQARRCLD+IDGAS+VLYG
Sbjct: 121 "CSDQFTIALKCYPRKYEDAQEIVDLLIVNECVLIDFQPMLDQAARRCLDFTIDGASKVLYG 180

      Query: 164 SLQKVGSSMFLITPANVMVDIEEMNI PKTGQRTSFDFDMKRR 205
      SLQKVGSSM+LL P+NV V+IEM IP T Q+ FDFDMKRR
10  Sbjct: 181 SLQKVGSSMYLAPSNVSVNIEEMTI PHTTQDIGFDFDMKRR 222

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 72

- 15 A DNA sequence (GBSx0072) was identified in *S. agalactiae* <SEQ ID 235> which encodes the amino acid sequence <SEQ ID 236>. This protein is predicted to be YlmH. Analysis of this protein sequence reveals the following:

```

Possible site: 35

20  >>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.3956 (Affirmative) < succ>
25  bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AAC95444 GB:AF068901 YlmH [Streptococcus pneumoniae]
Identities = 101/255 (39%), Positives = 161/255 (62%)

30  Query: 6 IYQHFRPEXYAFIHKIDHIAQYVENTYSPITTEPLNPREFKILESJIERRGSHYTTGGY 65
      IYQHF E+ F+ K + VE++Y+ T F+NP + K+L+ + + G +SG++
Sbjct: 5 IYQHFSIHDRPFLDKGMWIKKVEDSYAPFLTPPINPHQEKLLKLIKTYGLACSSSGEF 64

35  Query: 66 FQTRYVKVITIAPEYQLDMADFNLSLRIKYNKAFNHITHAKIMGTILNLYGVGRSILGD 125
      +EYV+V++ P+Y+Q + +DF +SL EI Y+ KP HITHAKI+GT++N LG++R + GD
Sbjct: 65 VSSEYVRVLLYPDYPOPEFSDFBISLQEIYVSNKFEHITHAKILGTVINQLGIERKLFGD 124

40  Query: 126 ILVBEGCAQVLVDQMNHLVHSVTKIGTASVOLAEVPLSKILLTPKQDIQKITVIASSLR 185
      ILV+E AQ+++ Q + KIG V L E P + + + + ++L+ SS R
Sbjct: 125 ILVDEBQAQIMINQOFLLLFQDGLKKIGRI PVSLEERPFTKIDKLEQYRRLDLVSSFR 184

      Query: 186 LDKILATILIKISRTQSTKLIEADKVKVNYATVNRVSEQLVBGDLISVRGYGRFLTNHNLG 245
      LD +L+ +L+SR Q+ +LIE V+VNY V++ + GDLSIVR +GR L+ L+ G
45  Sbjct: 185 LDVLLSNVLKLSRNQANQLIEKKLVQVNYHVVDKSDYTVQVGDLISVRKGRRLRLDQKG 244

      Query: 246 LTINQKYKLEVDKMI 260
      TK +K K+ V ++
50  Sbjct: 245 QTIGKKKITVQLLL 259

```

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 237> which encodes the amino acid sequence <SEQ ID 238>. Analysis of this protein sequence reveals the following:

```

Possible site: 56

55  >>> Seems to have no N-terminal signal sequence
      INTEGRAL Likelihood = -0.69 Transmembrane 46 - 62 ( 46 - 62)

----- Final Results -----
      bacterial membrane --- Certainty=0.1277 (Affirmative) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

- >GP: AAC95444 GB: AF068901 YlrH [Streptococcus pneumoniae]
Identities = 110/257 (42%), Positives = 161/257 (61%)
- Query: 7 IYQHFGHEYPFFIDRMSDMINRVEDYILLEVTEFLNPREVMILKSLIALTLDMFVSTDY 66
IYQHF E+ PF+D+ + I +VED Y +T P+NP + +LK L L S++
Sbjct: 5 IYQHFSEIRPFIDRMSDMINRVEDYILLEVTEFLNPREVMILKSLIALTLDMFVSTDY 64
- Query: 67 YPSEYGRVIAFGYYDLQSDPQIALWEISYQAKPNQLTHSQILGTLINELGVKRNLPD 126
SEY EV++ P Y+ E SDP+I+L RI Y KP LTH++ILGT+IN+LG++R LPGD
Sbjct: 65 VSSEYVRVILYFDYFPQPEFSDPEISLQRIVSNKPFSLTHAKILGT+IN+LG++R LPGD 124
- Query: 127 VFVMSGYAGLMIKRELDDYPLGTITTKIAKTSVKLRVNFQDLIRSINDSQTLDLVSSFR 186
+ V+ AQ+MI ++ L F + KI + V L E P + I ++ + LD+ VSSFR
Sbjct: 125 ILVDEERAQIMINQQFILLPDGLKKGRIIPVSLERPFTEKIDKLEQVRELDSVSSFR 184
- Query: 187 LDGVVATILKSRQTQVIALIEANKIKVNYRVANKASDNLVIGMVSIRGHSRFTLLADNG 246
LD +++ +LK SR Q LIE ++VNY V +K+ + +GC++S+R GR LL D G
Sbjct: 185 LDVLSNVLKLSRNCANQLISKLLVQVNYHVVDKSDYTVQVGLISVRKGRLLQKNG 244
- Query: 247 VTKHGKQKITLSKMIH 263
TK K+KIT+ ++ K
Sbjct: 245 QTKKEKKITVQLLSK 261

An alignment of the GAS and GBS proteins is shown below:

- Identities = 123/256 (48%), Positives = 177/256 (69%)
- Query: 6 IYQHFPEEYAFIHKIDHLAQVENTYSPFITTEFLNPREFKILESULVERGSHYTSQY 65
IYQHF SEY FI ++ + VE+ Y TEFNPRE IL+S++ + S Y
Sbjct: 7 IYQHFQHEYPFFIDRMSDMINRVEDYILLEVTEFLNPREVMILKSLIALTLDMFVSTDY 66
- Query: 66 FQTEYVGVIIAPEYVQLMDADFNLSLIEIKYNAKPNHLTHAKINGTLAYLGVKRSILGD 125
+ +EY +VIIAP YY L+ +DF ++L+EI Y AKFN LTH++I+GTL+N LGVR++ GD
Sbjct: 67 YPSEYGRVIAFGYYDLQSDPQIALWEISYQAKPNQLTHSQILGTLINELGVKRNLPD 126
- Query: 126 ILVDEGCAQVLNDSQWNLVHVSFTKIGTASVQLAEVPLSKLLTPKQDIQKLTVIASSLR 185
+ VE G AQ+++ ++ ++ + +TKI SV+L EV +L+ + Q L ++ SR R
Sbjct: 127 VFVMSGYAGLMIKRELDDYPLGTITTKIAKTSVKLRVNFQDLIRSINDSQTLDLVSSFR 186
- Query: 186 LDKILATILKISRTOSTKLEADKVKVNYATVNVSBQLVGDLISVRGYSRFTLLANLG 245
LD ++ATILK SRQT LIEA+K+KUNY N+ S+ LV GD++S+RG+GRFTL + G
Sbjct: 187 LDGVVATILKSRQTQVIALIEANKIKVNYRVANKASDNLVIGMVSIRGHSRFTLLADNG 246
- Query: 246 LTKHGKQKITLSKMIH 261
+TK+ K K+ + KMIH
Sbjct: 247 VTKHGKQKITLSKMIH 262

- Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 73

A DNA sequence (GBSx0073) was identified in *S. agalactiae* <SEQ ID 239> which encodes the amino acid sequence <SEQ ID 240>. This protein is predicted to be cell division protein DivIVA (septumplacement).

- Analysis of this protein sequence reveals the following:

Possible site: 14

>>> Seems to have no N-terminal signal sequence

- Final Results -----

bacterial cytoplasm --- Certainty=0.5418 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

5 The protein has homology with the following sequences in the GENPEPT database:

>GP: AAC95445 GB: AF068901 cell division protein DivIVA (Streptococcus pneumoniae)
 Identities = 132/227 (58%), Positives = 179/227 (78%), Gaps = 2/227 (0%)

10 Query: 1 MPTALRLKDKTFFSKFRGYSEEVNEFLITVDDYEDLIRNRBQEQYIKDLSEKIANF 60
 NP+T+LEIKDKTF ++FRG+ EEV+EFL+IVV DYEDL+R N ++ IK LBE++YF
 Sbjct: 1 NPITSLKDKTFTGTRFGFPOPEVEFDLIVDDYEDLVRANBDKNIRIKSLSEKLSYF 60

15 Query: 61 NEMKESLQGSVILAQETAEKRVKISQDASNLGKATFDAQHLIDEAKLANQILRDATD 120
 +E+K+SLQGSV++AQ+TAERVK +A ++N++ +A DAQ L++EAK KAN+ILR ATD
 Sbjct: 61 DEIKDSLQGSVILAQDTAEKRVQAHERSNWIIHQASQDQQLLEAKYKANEILRQATD 120

20 Query: 121 DAKRVAIETFDLKRQSRVPHQRLLESELEQLKANSAMWELKPTATYIQNSDAPKEV 180
 +AK+VA+ETE+LK +SRVPHQRL S +E QL + SS WE++L+PTA YLQ SD +PEV
 Sbjct: 121 NAKKVAVETKELKNSRVPHQRLKSTSSQSLAIVSSDWEDILRPTATYIQTSDAPKEV 180

25 Query: 181 VEKVLDEDDALPVVDITSPDATRQPSDENELQRRVRESNQLE 227
 V +VL E P+ + E D TRQPS EN ELQ R+E +K+L E
 Sbjct: 181 VSEVLGEPIPAPI--KEEPIDMTRQPSQAEMLQARIEVADKLE 225

25 A related DNA sequence was identified in *S. pyogenes* <SEQ ID 241> which encodes the amino acid sequence <SEQ ID 242>. Analysis of this protein sequence reveals the following:

Possible site: 14

>>> Seems to have no N-terminal signal sequence

30 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.6272 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

35 An alignment of the GAS and GBS proteins is shown below:

Identities = 180/254 (70%), Positives = 217/254 (84%), Gaps = 2/254 (0%)

40 Query: 1 MPTALRLKDKTFFSKFRGYSEEVNEFLITVDDYEDLIRNRBQEQYIKDLSEKIANF 60
 M LT LEIKDKTF +KFRGY SEEVNEFLITVDDYED L+RNR+ E IKDLSEK++YF
 Sbjct: 1 MALTTLEIKDKTFFTKFRGYCEEVNEFLITVDDYEDLVRANBDKNIRIKSLSEKLSYF 60

45 Query: 61 NEMKESLQGSVILAQETAEKRVKISQDASNLGKATFDAQHLIDEAKLANQILRDATD 120
 +EMKESLQGSVILAQETAE+K +A EA+NL+ KAT+DAQHL+DE+K KANQ+LRDATD
 Sbjct: 61 DEMKESLQGSVILAQETAEKRVKATANAATNLVSKATYDAQHLIDESKAKANQILRDATD 120

50 Query: 121 DAKRVAIETFDLKRQSRVPHQRLLESELEQLKANSAMWELKPTATYIQNSDAPKEV 180
 +AKRVAIETE+LKRQ+RVPHQRL+S +E QL L+NS W+ELL+PTATYIQNSDAPKEV
 Sbjct: 121 EAKRVAIETELKQRTVPHQRLISSISQSLSNSPWDELLQPTATYIQNSDAPKEV 180

55 Query: 181 VEKVLDEDDALPVVDITSPDATRQPSDENELQRRVRESNQLE 240
 V+ VL+ED +P DD+ SFDATRQPS+P+E+ELLQRRV+ESNK+LE L ++ E
 Sbjct: 181 VKTVLNE--IPESDSDSDATRQPTPKELKELQRRVDRSNKLELQYLDQSDSTTEP 238

60 Query: 241 PINLGETCTFKINI 254
 +NL ETCTFKINI
 Sbjct: 239 KVNLSCTCTFKINI 252

60 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 74

A DNA sequence (GBSx0074) was identified in *S.agalactiae* <SEQ ID 243> which encodes the amino acid sequence <SEQ ID 244>. Analysis of this protein sequence reveals the following:

Possible site: 61

>>> Seems to have no N-terminal signal sequence

INTEGRAL Likelihood = -0.43 Transmembrane 841 - 857 (841 - 857)

----- Final Results -----

bacterial membrane --- Certainty=0.1171(Affirmative) < succ>

bacterial outside --- Certainty=0.0000(Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAC95446 GB:AF068901 isoleucine-tRNA synthetase [Streptococcus pneumoniae]
Identities = 730/929 (78%), Positives = 822/929 (87%), Gaps = 1/929 (0%)

Query: 1 MKLKETLNLGQTAFPMRAGLPKPEQWQEWADQADLYKKRQALNKGKPAFLHLDGPPYAN 60

MLKk+TLNLG+T FPMRAGLP KEP WQ+ W+ A +Y++RQ LN+GKPF LHDGPPYAN

Sbjct: 1 MKLKDTLNLGKTEFFPMRAGLPTKEPFWQKEWDAKLYQRQELNQGKPHFTLHDGPPYAN 60

Query: 61 GNTHVGHALNKISKDIIVRSKSMGFPAPFYFGNDTHGLPIBQVLAKKGKVRKEMDLAEY 120

GNTHVGHkA+NKISKDIIVRSKSMGFP AP++FGNDTHGLPIBQVL+K+GVRKEMDL EY

Sbjct: 61 GNTHVGHkA+NKISKDIIVRSKSMGFPAPFYFGNDTHGLPIBQVLKSGKVRKEMDLAEY 120

Query: 121 LEMCRDYALSGVDKRDQDFKRLGVSADWENPYITLTDPYEAQKRVFGAMADKSYIYRGA 180

L++CR+YALSGVDKQR+DFKRLGVS DWENPY+TLTDPYEA Q+RVFG Mh+KSYIYRGA

Sbjct: 121 LKLCREYALSGVDKQREDQDFKRLGVSADWENPYITLTDPYEAQKRVFGMANKSYIYRGA 180

Query: 181 KPVYVWSSSBSALAEAEIEYHLDISTLYYANKVKDGKGLDITDTYIVVMTTPTFTVAS 240

KPVYVWSSSBSALAEAEIEYH+D STSLYANKVKDGKGLDITDTYIVVMTTPTFT+VAS

Sbjct: 181 KPVYVWSSSBSALAEAEIEYHLDVSTSLYANKVKDGKGLDITDTYIVVMTTPTFTVAS 240

Query: 241 RGLTVGDPMEYVVVVVFGSERKYLAEVLVDSLAAKPGWENFVITHTTGKELNHIVTER 300

RGLTVG D++YV+V FVG RK++A L+ SL+ KFGW + +++ + G+ELNHIVTER

Sbjct: 241 RGLTVGADIDYVLVQVPGEARKFVVAEELLTSLSEKPGWADQVLETRYQELNHIVTER 300

Query: 301 FWDTEVEELVILGDHVTDSGTGIVHTAPGFGEDDYNGIANGLDVVTVDSRGLANENA 360

FWDTEVEELVILGDHVTDSGTGIVHTAPGFGEDDYNGIANG L+V VTVD R+NN+NA

Sbjct: 301 FWDTEVEELVILGDHVTDSGTGIVHTAPGFGEDDYNGIANGLEVAVTVDSRGINNNNA 360

Query: 361 GPDFEGQGFYDKVTPLVKKEKGLDLLSLRVINSHSYDFWRTKKPIINRAVQPMFASVSKR 420

GP+PBGQGFY+KV P V EKLG+LLLA I B+HSYPDFWRTKKPIINRAVQPMFASVSKR

Sbjct: 361 GPDFEGQGFYKVVPTVIEKGLNLLAQESISHSYDFWRTKKPIINRAVQPMFASVSKR 420

Query: 421 QEILDEIKETNPQPMKSKRLRYNMIRNDRGDWISQRQAWGVLPPIFYAEDGTAIMTKETV 480

QEILDEIK E F EMKG RLYNMIRNDRGDWISQRQ WGVLPPIFYAEDGTAIM E

Sbjct: 421 QEILDEIKETNPQPMKSKRLRYNMIRNDRGDWISQRQWGVLPPIFYAEDGTAIMVASTI 480

Query: 481 DHVADLFASYSIGSIWQRIAKOLLEPAGYTHPGSPNGFKEKSTIDIMVWFDSSGSSWGVN 540

+HVA LF ++GS +W+RDAKOLLE +GTHPGSPNG F+KSTIDIMVWFDSSGSSWGVN

Sbjct: 481 EHVADLFKKGSSIIWRIAKOLLEPAGYTHPGSPNGFKEKSTIDIMVWFDSSGSSWGVV 540

Query: 541 ARENLSPADLYLBSGDYQWGFNSSLITSVANNGHAPYKAVLSQGFVLGDKGKMSKSL 600

R L+YPADLYLBSGDYQWGFNSSLITSA +G APYK +LQGF LDGDKGKMSKSL

Sbjct: 541 NRPELITPADLYLBSGDYQWGFNSSLITSVANNGHAPYKQILSQGFALDGGKMSKSL 600

Query: 601 GNTILPSDVEKQGAELRLWVTSVDSNDVRISMDILKQSETYRKIRNTLRFLIANTS 660

GNTI PSDVEKQGAELRLWVTSVDSNDVRISMDIL Q SETYRKIRNTLRFLIANTS

Sbjct: 601 GNTILPSDVEKQGAELRLWVTSVDSNDVRISMDILSQVSETYRKIRNTLRFLIANTS 660

Query: 661 DFNPKQDAVAYENLGAVDYMTIKFQVVTINKAYADFMAYIKAVNMFVTLDSAFY 720

DFNP QD VAY L +VD+YMTI+FNQ+V TI AYA ++P+ IYKA+VNP+ VDLDSAFY

Sbjct: 661 DFNPAQCTVAYDELASVVKYMTIRFQVLTIRDAYDFEFLIYKALWNTINVLDSAFY 720

-138-

Query: 721 LDFAKDVVYIEANSPERRRMQTVFYDILVKLTLL/TPILPHTAEIRWSYLESHHEEEFVQ 780
 LDFAKDVVYIE A S ERR+MQTVFYDILVK+TKLL/TPILPHTAEIRWSYLE E B+PVC
 5 Sbjet: 721 LDFAKDVVYIEGAKSLERQMQTVFYDILVKLTLL/TPILPHTAEIRWSYLEFEFEDFVQ 780

Query: 781 LAEMPVQAQTFSGQSEILSEWSAFMTLQTAQQALEARNNAKVIGKSLAHLTIYASQEVK 840
 L+E+P QTF+ QSEIL+ N+AFM R QQAQALEARNNAKVIGKSLAHLT+Y + + VK
 Sbjet: 781 LSELPEVQTFANQSEILDTWAAFMDFPQQAQQALEARNNAKVIGKSLAHLTVYENEVK 840

10 Query: 841 TLLTALNSDIALLMIVSQLTIADEADKPADSVSFEGVAFVIEHAGRCVRSRRIDPTTK 900
 TLL A+NS++A L+IVS+LTIA+E P ++SFE VAFVIE A GEVC+R RRIDPTT
 Sbjet: 841 TLLAEVNSVAGLLIVSELTIAEE-PAPEAALSFEVAFVIEAAGVCDCRCRIDPTTA 899

15 Query: 901 MRSYGVAVCDASAAIIBQYYPEAVAGQFE 929
 RSY +CD A+I+E+ + +AVA+GFE
 Sbjet: 900 ERSYQAVICDHCAISIVEKNFADAVAEGFE 928

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 245> which encodes the amino acid sequence <SEQ ID 246>. Analysis of this protein sequence reveals the following:

20 Possible site: 61

>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -1.70 Transmembrane 849 - 865 (848 - 867)

25 ----- Final Results -----
 bacterial membrane --- Certainty=0.1680 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

30 An alignment of the GAS and GBS proteins is shown below:
 Identities = 798/929 (85%), Positives = 857/929 (91%)

Query: 1 MKLKETLNLGQTAPPMRAGLPNKEPQWQAEWQADIIYKKRQALNEGKPAFHLHDGPPYAN 60
 MKLKETLNLG+TAPPMRAGLPNKEPQWQ AM+QA++YKKRQ IN GKPAFHLHDGPPYAN
 35 Sbjet: 1 MKLKETLNLGQTAPPMRAGLPNKEPQWQAAWEQAEIYKKRQELNAGKPAFHLHDGPPYAN 60

Query: 61 GNIHVGHALNKISKDIIVRSKSMSCGFAPYVFGWDTGHLPIEQVLAKKGKRRKMDLAEY 120
 GNIHVGHALNKISKDIIVRSKSMSCGF+APYVFGWDTGHLPIEQVLAK+G+KRRKMDLAEY
 Sbjet: 61 GNIHVGHALNKISKDIIVRSKSMSCGFAPYVFGWDTGHLPIEQVLAKGGIKRRKMDLAEY 120

40 Query: 121 LEMCRDYLALSOVDKQRDDFKRLGVSADWENPYITLTPDYEADQVRVFGAMADKGYIYRGA 180
 LEMCR YALGQVDKQRDDFKRLGVSADWENPY+TL P +RADQ+RVFGAMA+KGYIYRGA
 Sbjet: 121 LEMCRQYALSOVDKQRDDFKRLGVSADWENPVITLDPQFEADQIRVFGAMAEKGYIYRGA 180

Query: 181 KPVIYWSSESALAEAEIEYHDIDSTSLYANKVKDGKILDTITVIYVWTTTPTPTVAS 240
 KPVIYWSSESALAEAEIEYHDIDSTSLYANKVKDGKILDT+TVIYVWTTTPTPTVAS
 Sbjet: 181 KPVIYWSSESALAEAEIEYHDIDSTSLYANKVKDGKILDTNTVIYVWTTTPTPTVAS 240

50 Query: 241 RGLTVGPDMEYVVVVPVSGSERKYLAEVLVDSIAKPGOWENFEIVTHHTQKLNHIVTEH 300
 RGLTVGPDW+Y+VV P GS+R+Y++AE L+DSL A KPGW+FR + H G +L +IVTIEH
 Sbjet: 241 RGLTVGPDMDYLVPKPGSDRQYVVGAGLLSLAGKPGWSEFETLASHKGADLEIVTIEH 300

Query: 301 FWDITVEEELVILGDHVTLSGIGIVHTAPGFGEDDYNVGIANGLDVVVTVDSRGLMMENA 360
 FWDIT+VEEELVILGDHVT +SGVIGIVHTAPGFGEDDYNVG L+V VTVD RGLMMENA
 55 Sbjet: 301 FWDITVEEELVILGDHVTLESIGIGIVHTAPGFGEDDYNVGTIKLEVAVTVDSRGLMMENA 360

Query: 361 GPDFGQGFYQKVTPLVKEKIGDLLLAGEVINSYFPDWRKKPLIIRAVPQWASVSQKFR 420
 GPDF GQFY+KVT+V +KIGDLLLA EVINSYFPDWRKKPLIIRAVPQWASVS FR
 Sbjet: 361 GPDFHGQGFYQKVTPIVDKIGDLLLAGEVINSYFPDWRKKPLIIRAVPQWASVSDFR 420

60 Query: 421 QEILDEIEKTNPQBPBGKKRLYNNMIRDRGDWVISRQAWGVPLPIFYAEDGALMTKEVT 480
 Q+ILDEIEKT P P WG+ KLYNNMIRDRGDWVISRQAWGVPLPIFYAEDGALMTKEVT
 Sbjet: 421 QDILDEIEKTT+PHPSWGETRLYNNMIRDRGDWVISRQAWGVPLPIFYAEDGALMTKEVT 480

65 Query: 481 DEIVADLAEYGSIVWQRDAKDLIPAGYTHFGSPNGLPEKETDINDVWFDPSQSNWGVN 540

```

      DHVADLF E GSI+WWQ++AKDLLF G+THPGSPNG F KETDIDMVNFDGSSGWNQVMN
Sbjct: 481 DHVADLFQENGSIINWQKEAKDLLFEGPTHGSPNGEPTKETDIDMVNFDGSSGWNQVMN 540
5  Query: 541 ARENLSYPADLYLKGSDQYRGWFNSSLITSAVNGHAPYKAVLSQGFVLGKGKRMKSL 600
      +ENLSYPADLYLKGSDQYRGWFNSSLITSAVNGHAPYKA+LSQGFVLGKGKRMKSL
Sbjct: 541 TKENLSYPADLYLKGSDQYRGWFNSSLITSAVNGHAPYKAVLSQGFVLGKGKRMKSK 600
      GNTILPSDVEKQPGARILRLMVTSDSSNDVRISMILKQTSYRKIRNTLRLFLIANTS
10 Sbjct: 601 GNTILPSDVEKQPGARILRLMVTSDSSNDVRISMILKQTSYRKIRNTLRLFLIANTS 660
      GN I P+DV KQ+GA+ILRLMV SVD+ NVR+SM+IL Q SETYRKIRNTLRLFLIANTS
Query: 661 DFNPKQDAVAYENLGAVDYMTIKFNQVVTINKAYAAYDFMAIKAVNVFVTLDSAFY 720
      DFNPK D VAY +LG VD+YMTI FNQ+V TI AY YDFMAIKAVNVFVTLDSAFY
15 Sbjct: 661 DFNPKDPTVAYADLGTVDKYMFTVFNQLVATITDAYERYDFMAIKAVNVFVTLDSAFY 720
      LDFAKDVVYIEAANSFERRRMQTVFYDILVKITKLTLPILPHTAERIMSYLEHEEESFVQ
Query: 721 LDFAKDVVYIEAANSFERRRMQTVFYDILVKITKLTLPILPHTAERIMSYLEHEEESFVQ 780
      LDFAKDVVYIEAANS FERRRMQTVFYDILVK+TKLTLTPILPHT EIMSYLEHE E FVQ
Sbjct: 721 LDFAKDVVYIEAANSFERRRMQTVFYDILVKITKLTLPILPHTIEIMSYLRHESEAFVQ 780
20 Query: 781 LAEMFVACFTSGQRILEKWSAPMTLKTQAKALEEARNAKVIGKSLAHLTIYASQEVK 840
      LAEMFVA+TFS QE+ILS WSAFMTLKTQAKALEEARNAK+IGKSLAHLTIYAS+EVK
Sbjct: 781 LAEMFVARTFSAQSDILKAWSAFMTLKTQAKALEEARNAKIIGKSLAHLTIYASQEVK 840
      TLLTALNSDIALIMIVSQLTIADEADKPADSVSFEQVAFVHEAHEGVCERSRRIDPTTK
25 Query: 841 TLLTALNSDIALIMIVSQLTIADEADKPADSVSFEQVAFVHEAHEGVCERSRRIDPTTK 900
      TLLTAL+SDIALI+IVSQLTIAAD PAD+V+FEQVAF VEHAE GVCERSRRIDPTT+
Sbjct: 841 TLLTALNSDIALI+IVSQLTIAADAPADAFAFEQVAFVHEAHEGVCERSRRIDPTTR 900
      MRSYGVAVCDASNAALIEQYYPEAVAGGFE 929
Query: 901 MRSYGVAVCDASNAALIEQYYPEAVAGGFE 929
      MRSY VCD SA IIE+ +PEAVA+GFE
30 Sbjct: 901 MRSYNAFVCDHSAKILIEENFPEAVAEGFE 929

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 75

35 A DNA sequence (GBSx0075) was identified in *S. agalactiae* <SEQ ID 247> which encodes the amino acid sequence <SEQ ID 248>. Analysis of this protein sequence reveals the following:

```

Possible site: 39
>>> Seems to have no N-terminal signal sequence
---- Final Results ----
      bacterial cytoplasm --- Certainty=0.3425 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
45      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 249> which encodes the amino acid sequence <SEQ ID 250>. Analysis of this protein sequence reveals the following:

```

Possible site: 32
>>> Seems to have no N-terminal signal sequence
---- Final Results ----
      bacterial cytoplasm --- Certainty=0.3467 (Affirmative) < succ>
55      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

Identities = 77/99 (77%), Positives = 89/99 (89%)

Query: 1 MRLINTTSSHPVLVRNQLQNTDAKLVEVYSAGNTDVVFTKAPKHYELLISNKYRAIKDEE 60
 MRLINTTSSHPVL++NQL+NTDA LVEVYSAGNTDV+FT+APKHYELLISNKYRAIK++E
 Sbjct: 1 MRLINTTSSHPVLKLNQLQNTDAYLVEVYSAGNTDVVFTKAPKHYELLISNKYRAIKDEE 60

Query: 61 LEAIREFPLKRRKIDQSIILQBMKSLWPAKLIRISYPT 99
 L+ IREFFLKRRKID I+I Q K+LHT LIEIS+ T+
 Sbjct: 61 LDIREFPLKRRKIDPKIVIPQGSKTLWNNLIRISPTS 99

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 76

A DNA sequence (GBS0076) was identified in *S. agalactiae* <SEQ ID 251> which encodes the amino acid sequence <SEQ ID 252>. This protein is predicted to be AP4A hydrolase. Analysis of this protein sequence reveals the following:

Possible site: 42

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1714 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAC06510 GB:AE000676 AP4A hydrolase [Aquifex aeolicus]
 Identities = 30/101 (29%), Positives = 49/101 (46%), Gaps = 2/101 (1%)

Query: 32 KILVQAPNGAWFLPGGEIEENNNHLSALTRELIELGYSATIGHYQGADEYFYSRHRD 91
 +++L++ P+ W P G I E E E RE+ EE G I Y G+ Y+Y+ +
 Sbjct: 16 EVLLIKTPGNVNSPFPKGNIEPCKPEETAIVREVWEETGVKGELIDYIGEI-HYNYTLKGE 74

Query: 92 TYYNPAIYIEVTAYHKDQAPLEDNFHLAWFFIQEAKELK 132
 + Y Y + + P + +PFI+RAK+ LK
 Sbjct: 75 RIFKTVKY-VLMKYKSGEPRPSWEVKDAKFFFIKERKLLK 114

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 253> which encodes the amino acid sequence <SEQ ID 254>. Analysis of this protein sequence reveals the following:

Possible site: 47

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1954 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 102/149 (68%), Positives = 118/149 (78%)

Query: 1 MTNPTFCEKIDNNYRSRFGVYAIIPNPTDKILVQAPNGAWFLPGGEIEENNNHLSAL 60
 M PTPG K + +Y +R+GVYAIIPN KILVQAPNG+WFPLPGGEIR E L+L+
 Sbjct: 1 MMLPTFGHKNHNDYVTRYGVYAIIPNHBQTKILVQAPNGSWFLPGGEIRAGSGQLQAL 60

Query: 61 RELIELGYSATIGHYQGADEYFYSRHRDTYYNPAIYIEVTAYHKDQAPLEDNFHLA 120
 RELIELG+SATIG YTGGADEYFYSRHRDT++Y+PAY+YRVT+ PLEDNF+L
 Sbjct: 61 ERELIELGYSATIGSYGGADEYFYSRHRDTHFYHPAYLVEVTAQAVSKPLEDFNFG 120

Query: 121 WFFIQEAKEKLGRGSHRWGVQVAMEKNHHS 149
 WF EA KLGR SH+WG+V W+K HHS
 Sbjct: 121 WFSPTIEAIAKLGRSHRWGVQVAMEKNHHS 149

- 5 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 77

A DNA sequence (GBSx0077) was identified in *S. galactiae* <SEQ ID 255> which encodes the amino acid sequence <SEQ ID 256>. This protein is predicted to be ClpE (clpB-1). Analysis of this protein sequence
 10 reveals the following:

Possible site: 54

>>> Seems to have no N-terminal signal sequence

- 15 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.2882 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

- 20 The protein has homology with the following sequences in the GENPEPT database:

>GP:AAD01782 GB:AF023421 ClpE [Lactococcus lactis]
 Identities = 560/752 (74%), Positives = 647/752 (85%), Gaps = 12/752 (1%)

- Query: 1 MLCQNCKLNESITHLTYTNVNGKQKQVLDQCNCYQIKTDPNNPLFSGNLNHS-HAFGGIN 59
 25 MLCQNC +NE+THLTY+VNG++K+DLCQNCYQI+K+ LF N + ++ N
 Sbjct: 1 MLCQNCNINEATILHTSVYNGKKQIDLCQNCYQIMKSGGQALFAGNANSGNSDEPFN 60

- Query: 60 PFFDDFFGDLNNFAPNGQDLPTPTTQSGGNRGCGNNGRNNRNNOTATPSQAKGILEE 119
 30 PF +D F L + PNG TPPTQ+GG G NR Q KG+LEE
 Sbjct: 61 PF-NDIFPSALQG-QDPPGAASNGTPTTQSGRGRPRGQNPFR-----AKQPKGNLEE 109

- Query: 120 FGINVTEIARHGIDIPVIGRDSIIRVIBILNRRTKNNFVLIGEPGVGKTA VVEGLAQKI 179
 FGIN+TE AR G+IDPVIGR DI RVIRILNRRTKNNFVLIGEPGVGKTA VVEGLAQKI
 Sbjct: 110 FGINITESARRGEIDPVIGRDEIRKRVIBILNRRTKNNFVLIGEPGVGKTA VVEGLAQKI 169

- 35 Query: 180 VDGNVPHKLQKQVIRLDVSVLQGTGIRGQFEBRMQKLMSEIRQRQDVILFIDEIHEIV 239
 VDG+G+P KLQ K+VIRLDVSVLQGTGIRGQFEBRMQKLM+EIR+R DVI+FIDEIHEIV
 Sbjct: 170 VDGCVPKLQKQVIRLDVSVLQGTGIRGQFEBRMQKLMDEIRKRNVDVIMFIDEIHEIV 229

- 40 Query: 240 GAGTAGGSGVDAGNLLKPALARGELQLVGATTILNEYRIIEKDAALESRRMQPVKVDPSVE 299
 GAG+G+G+YDAGNLLKPALARGELQLVGATTILNEYRIIEKDAALESRRMQPVKVDPSV+
 Sbjct: 230 GAGSAGDGNVDAGNLLKPALARGELQLVGATTILNEYRIIEKDAALESRRMQPVKVDPSVD 289

- Query: 300 ETITILGIGIKKYEDYHIVKYNNDAIEPAAVLNRYIQDRFLPKAIDLLDEAGSRNNLT 359
 45 ETITIL+GIGI+YEDYHIVKY ++AIRAA LSNRYIQDRFLPKAIDLLDE+GSK NLT
 Sbjct: 290 ETITILRGIGIARYEDYHIVKYTDIEAPAAVLNRYIQDRFLPKAIDLLDEAGSRNNLT 349

- Query: 360 LNFVDPKEIDQRLIEARNLKAQATREEDYERAAVPRDQIAKYKEMQQQKVDQDPTIITE 419
 50 L FVDP++I++R+ +AE+ K +AT+ ED+EA++PRDQI+K +E+Q+Q+V D+D P+ITE
 Sbjct: 350 LKFDVPEIDNRIADAEKNEATKAEDPEKAAHPRDQISKLELQKQEVTDMDP+ITE 409

- Query: 420 KTIIRHIEBKTNIPVDLKRKESQLINLADLLKHVIGQDDAVVAKIARRNRVGLGS 479
 K IE I+E+K+T IPVDLKRKQ+QLNINLADLL KHVIGQD+AV KI+K+IR+R+RVGLG
 Sbjct: 410 KDIQIVQEQTQIPVDLKRKESQLINLADLLKHVIGQDRAVDKISKARRSRVGLQK 469

- 55 Query: 480 PNRPIGSLFVPGFTGVGKTELKQLAIELPGSADSMIRFDNSEYMEKHAVAKLVGAPPOY 539
 PNRPIG FLVPGFTGVGKTEL+KQLA ELFGS++SMIRFDNSEYMEKH+VAKL+GAPPGY
 Sbjct: 470 PNRPIGFLFVPGFTGVGKTELAKQLAKELFGSS+SMIRFDNSEYMEKHVAKL+GAPPGY 529

- 60 Query: 540 VGYEAGQLTEKVRNNPYSILLLDEIRKAHFDVMRMFLQLVLDORLITDQGRVTSFKDITI 599
 VGYEAGQLTE+VRNNPYSILLLDEIRKAHFDVMRMFLQ+L+DGRITD QGRVTSFKD++

Sbjct: 530 VGYEAAQLQTERVRENFYSLILLDEIEKAHPDVMEFMFLQILSDGRLTDAQGRTVSFKDSL 589

Query: 600 IINTSNAGSGKTEASVGPASRGRTINSVLGLGNFFSPENNRFDGIIIFKALDKENLL 659

IINTSNAG+GK EASVGPGA+RGRT SVLGLGL+FFSPENNRFDGIIIF AL KENLL

Sbjct: 590 IINTSNAGTGKVEASVGPAAKRGKTSVLGLGDFSPENNRFDGIIIFKALSKEKLL 649

Query: 660 NIVDIMLSDVNARLAINGIHLDVDKVKKEIADVLGYDPMGARPLRRTIQSHIDATIDY 719

IVD+ML +VN ++ N IHL VT KEKIVDLGY+P MGARPLRR IQE+IED+I D+

Sbjct: 650 KIVDIMLDEVNBQIGRNDIHLSVTQAQKKEIADVLGYDPMGARPLRRIQENIEDSDADF 709

Query: 720 YLENFSEKELRAIMTSNGNIISKSKKTEEST 751

Y+E+P K+L A + + +I +++T B+T

Sbjct: 710 YIEHFETKQLVADLLDDKIVISNQTQBTART 741

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 257> which encodes the amino acid sequence <SEQ ID 258>. Analysis of this protein sequence reveals the following:

Possible site: 43

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3104 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 640/751 (85%), Positives = 691/751 (91%), Gaps = 7/751 (0%)

Query: 1 MLCQNCKLNBSITHLYTVNGKQKQVDLCQNCYQIIKDPPNPLFSGLNHVSHAPG-GIN 59

MLCQNC LNBSITHLYT+VNGKQ+QVDLCQNCYQI+K+DP N + +GL A +

Sbjct: 1 MLCQNCKLNBSITHLYTSVNGKQKQVDLCQNCYQIMKSDPANSILNGITPGYRAQDRGST 60

Query: 60 PFFDDFPGLNPFRAFNGQDLNPTPTQSGNGGNGNGNRNNRQTATPS---QAKG 115

PFFDDFPGLNPFRAF +LNTPTPTQ+G N GG G N N + A P QAKG

Sbjct: 61 PFFDDFPGLNPFRAF--NLNPTPTQAGQNGNGGGRVYGNVNGRPAQPTQNGQAKG 118

Query: 116 ILBEPGINVTIARHGIDIPVIGRDSBIIRVIEILNRRTQNNPVLIGEPGVKTA+VBSGL 175

+LBEPGINVT+IAR+G+IDPVIGRD EI RVIEILNRRTQNNPVLIGEPGVKTA+VBSGL

Sbjct: 119 ILBEPGINVTDIARNGNIDPVIGRDEBITRVIEILNRRTQNNPVLIGEPGVKTA+VBSGL 178

Query: 176 AQKIVDGNVPHKLQKQKQVIRLDVVSIVQGTGIRGQFEERMQKLMBSIRORQDVLFDIEI 235

AQKI+DG VP KLQKQKQVIRLDVVSIVQGTGIRGQFEERMQKLMBSIR R+DVLFDIEI

Sbjct: 179 AQKIIDGTVPQKLQKQKQVIRLDVVSIVQGTGIRGQFEERMQKLMBSIRNRKDVLFDFIEI 236

Query: 236 HEITVGAGTAGBGSDMCAENILKALARGELQLVGATTINBYRIIEKDAALRRMQPVKVD 295

HEITVGAG+AG+G+MDAGNLIKALARGELQLVGATTINBYRIIEKDAALRRMQPVKVD

Sbjct: 239 HEITVGAGSAGDGNMDCAENILKALARGELQLVGATTINBYRIIEKDAALRRMQPVKVD 298

Query: 296 PSVESTITILKGIGIKKYEDYHHVKYNDIAEAAVLGNRYIQDRFLPDKAIDLDDIAGSK 355

PSVESTITILKGIGI KYEDYHHVKY+ ATRAA LSNRYIQDRFLPDKAIDLDDIAGSK

Sbjct: 299 PSVESTITILKGIGIKKYEDYHHVKYSPAAIRAAHLGNRYIQDRFLPDKAIDLDDIAGSK 358

Query: 356 MNLTILNFVDPKIEDQRLIEAENLKAQATREDEYRAAYFRDQITAKYKMQQQKVDQDTP 415

MNLTILNFVDPKED+KLIEAENLKAQATREDEYRAAYFRQI KYK39Q QVD+QD P

Sbjct: 359 MNLTILNFVDPKIEDKRLIEAENLKAQATREDEYRAAYFRDQITKYK39QKVDQDTP 418

Query: 416 IITTEKTIEHIEKKTNI PVGDLKKEKBSQLINLADDLKQHVIGQDDAVVIAKAIKRRNRV 475

IITTEKTIE I+E+KTNI PVGDLKKEKBSQL+NL+A+DLK HVIGQDDAV KIAKAIKRRNRV

Sbjct: 419 IITTEKTIEAIVKQKTNI PVGDLKKEKBSQLVNLANDLKAHVIGQDDAVDKIAKAIKRRNRV 478

Query: 476 GLGSPNRPIGSGFLPVGPTGVGKTELKSKQLAIELFSGSADMIRPDMSEYMEKHAVKLWGA 535

GLG+PNRPIGSGFLPVGPTGVGKTELKSKQLAIELFSGSADMIRPDMSEYMEKHAVKLWGA

Sbjct: 479 GLGTPNRPIGSGFLPVGPTGVGKTELKSKQLAIELFSGSTNMIRPDMSEYMEKHAVKLWGA 538

Query: 536 PFGYGVTEAQAQLTEKVRNPNFYSLILLDEIEKAHPDVMEFMFLQILDDGRLTDAQGRTVSF 595

```

PPGY+GYEAGQLTE+VRRNPYSLLILDE+EKAPDVMHMFQLVLDGRITDQGRITVSF
Sbjct: 539 PPGYIGYEAGQLTEQVRRNPYSLLILDEVRKAHPDVMHMFQLVLDGRITDQGRITVSF 598

Query: 596 KDTIIMTSNAGSGKTKASVGFASRSGRTNSVLGQLGNFFSPPEFNNRFDGLIEFKALDK 655
          KDTIIMTSNAG+GK+EASVGFGA+RBGRT+SVLG+L NFFSPPEFNNRFDGLIEFKAL K
Sbjct: 599 KDTIIMTSNAGTYSKSEASVGFGAARSGRTSSVLGSLNFFSPPEFNNRFDGLIEFKALSK 658

Query: 656 ENLNTIVDIMLSDVNARIADNGIHLDVTQKVEKIVDLSYDPKMGARPLERTIQERIEDA 715
          E+LL+IVD+ML DYN RL NQIHLDVT KVEKIVDLSYDPKMGARPLERTIQ+IESDA
Sbjct: 659 EHLLHIVDMLRDVNERLGYNGIHLDVTQKVEKIVDLSYDPKMGARPLERTIQEYIEDA 718

Query: 716 ITDYYLENPSEKKLRIMNTSNGNIIKSSKK 746
          ITDYYLE+P+EK+LRA+NT++ NI IK+ K+
Sbjct: 719 ITDYLEHPTKQLRALMNTSNKNTIKAVKE 749

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 78

A DNA sequence (GBSx0078) was identified in *S.galactiae* <SEQ ID 259> which encodes the amino acid sequence <SEQ ID 260>. This protein is predicted to be glutamine ABC transporter, permease protein (glnP). Analysis of this protein sequence reveals the following:

Possible site: 61

```

>>> Seems to have an uncleavable N-term signal seq
INTEGRAL Likelihood = -9.92 Transmembrane 27 - 43 ( 15 - 46 )
INTEGRAL Likelihood = -2.50 Transmembrane 200 - 216 ( 196 - 217 )

----- Final Results -----
          bacterial membrane --- Certainty=0.4970 (Affirmative) < succ>
          bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
          bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 9619> which encodes amino acid sequence <SEQ ID 9620> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AAB91000 GB:AB001090 glutamine ABC transporter, permease protein
(glnP) [Archaeoglobus fulgidus]
Identities = 92/209 (44%), Positives = 129/209 (61%), Gaps = 10/209 (4%)

Query: 17 YGVVMTIMISTCVVFGTIIGVLIALVKRTNLHFLTLANFYVWVFRGTPMVQIMIAFA 76
      +G VT+ ++ +FFG IIG + L + + ++ XV V KSTP++VQI+I +
Sbjct: 21 FGASVTLKLTLSIFPGLLIGTLAGLGRVSKNPLPFAISTAVEVRGRTPLVQILLIVF 80

Query: 77 WHHFNPLPTISFGVLDDLPFRLLPGIIIIISNGAYISRIVRAGIEAVPESQIRARYSLG 136
      LP I + GII +S+ SGAYI+EIVRAGIE++P GQ+RAA SLG
Sbjct: 81 ----GLPAIGINLQPEP-----AGIIALSICGAYIAKIVRAGISIPIGMRAARSIG 130

Query: 137 IRPKNTIRYIVLPQAFKNILPALGNRFITTIKDSALLQITIGVMELNNGASVW+ATYSPV 196
      + +RYVI PQAF+NLIPALGNRFI++KDS+LL I ++EL + +V T++
Sbjct: 131 NTYLQAMRYIVFPQAFKNILPALGNRFIALLKDSLSVISIVELTVRGQIVNTTNNAW 190

Query: 197 APLLFAFYFYLMLTLLSALLQMEKYIG 225
      P L A +YLM+T LS L+ +K LG
Sbjct: 191 TPPLGLVALFYLMITPLSLRWAYSQKLG 219

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 261> which encodes the amino acid sequence <SEQ ID 262>. Analysis of this protein sequence reveals the following:

Possible site: 30

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```
>>> Seems to have an uncleavable N-term signal seq
INTEGRAL    Likelihood = -9.08    Transmembrane    25 - 41 ( 11 - 44)
INTEGRAL    Likelihood = -1.91    Transmembrane    202 - 218 ( 201 - 218)

----- Final Results -----
bacterial membrane --- Certainty=0.4630(Affirmative) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>
```

The protein has homology with the following sequences in the databases:

```
>GP:AA891000 GB:AE001090 glutamine ABC transporter, permease protein
(glnP) [Archaeoglobus fulgidus]
Identities = 91/209 (43%), Positives = 138/209 (65%), Gaps = 12/209 (5%)

Query: 15 YGVLVTIMISVSVPFGTLIGVLVTLIKRSHVKPLIUVNL-VYVIFPGTGMVQIMIAF 73
+G VI+ + + + + PFG -IG + L + S PL + + + YV + RGP+ +UQI+I +
Sbjct: 21 FGASVTLKLTLISIPFGLIIGTLAGLRVSK-NPLPFAISTAYVEVIRGTPLLVQILIVY 79

Query: 74 ANNHFNMPITIGFGLDLDFSRLLPGIIISLNSGAYISEIVRAGIEAVPKGQLEAAVSL 133
+P IG + + + GII +S+ SGAYI+EIVRAGIE+P GQ+EA SL
Sbjct: 80 F-----GLPAIG-----INLQEPAGIIALSICSGAYIAEIVRAGIESIPGQMEAARSL 129

Query: 134 GIRPQAMRYVILPQAFKNILPALGNEFTTIKDSALLQTIGVMELNNGAQSVVITATYSP 193
G+ ANRYVI PQAF+NILPALGNEFTI ++KDS+LL I ++EL + +V T++
Sbjct: 130 GMTYLQAMRYVIFPQAFKNILPALGNEFTIILKDSLSVISIVELTVGRQIVNCTFMA 189

Query: 194 ISPLLVAAFYILMVTVMQALLAVLERHM 222
+P L A +YLM+T ++L+A ++ +
Sbjct: 190 WTPFLGVALFYLMVTIPLSRIVAYSQKKL 218
```

An alignment of the GAS and GBS proteins is shown below:

```
Identities = 180/225 (80%), Positives = 208/225 (92%)

Query: 3 MNFSFLPQWMSYFNGVMTIMISTCVVFFOTLIGVLIALVKPTILHFLTILANFYVWF 62
M+ SFLP+YW+YFNGV+VTIMIS VVFFOT+IGVL+ L+KR+++ LT + N YW+P
Sbjct: 1 MELSFLPKWYAFNGVULVTIMISVSVPFGTLIGVLVTLIKRSHVKPLIUVNLVYVWF 60

Query: 63 RGTPMVQIMIAFAMHGFNNLPTISFGVLDLDPTRLLPGIIISLNSGAYISEIVRAGIE 122
RGTPMVQIMIAFAMHGFNN+PTI FGVLDF+RLLPGIIISLNSGAYISEIVRAGIE
Sbjct: 61 RGTPMVQIMIAFAMHGFNNMPTIGFVLDLDFSRLLPGIIISLNSGAYISEIVRAGIE 120

Query: 123 AVPSGQIEAAVSLGIRPKVILRVILPQAFKNILPALGNEFTTIKDSALLQTIGVMELN 182
AVP GQ+EAAYSLGIRP+N +RVILPQAFKNILPALGNEFTTIKDSALLQTIGVMELN
Sbjct: 121 AVPKGLEAAVSLGIRPQAMRYVILPQAFKNILPALGNEFTTIKDSALLQTIGVMELN 180

Query: 183 NGAQSVVITATYSP+PILLFAAFYILMTTILSALLQMKRYLQKG 227
NGAQSVVITATYSP++PLL AAFYILMTT+++ LL +E+++ +G
Sbjct: 181 NGAQSVVITATYSPISPLVAAFYILMVTVMQALLAVLERHMAQG 225
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 79

A DNA sequence (GBSx0079) was identified in *S.galactiae* <SEQ ID 263> which encodes the amino acid sequence <SEQ ID 264>. This protein is predicted to be phosphomannomutase (manB). Analysis of this protein sequence reveals the following:

Possible site: 60

>>> Seems to have no N-terminal signal sequence

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----- Final Results -----

bacterial cytoplasm --- Certainty=0.5400 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

5

A related GBS nucleic acid sequence <SEQ ID 9621> which encodes amino acid sequence <SEQ ID 9622> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

10 >GP:BA004825 GB:AP001510 phosphomannomutase [Bacillus halodurans]
 Identities = 239/548 (43%), Positives = 344/548 (62%), Gaps = 14/548 (2%)

15 Query: 4 MNYKEI YQEWLENDLSIGKDIKSLEAIGDESEIQDRFYKTLFPGTAGLRGLGAGNRM 63
 M++++ Y++W + L ++K LEAI GDE +++D FYK LEFGT G+RG+G G NRM
 Sbjct: 1 MSWRQRYSKWKGFNELESELEKQSLGAIGGDEQLEDCFYKNLEFGTGMGRGSGIPGNRM 60

20 Query: 64 NTYMWGKAAQALANTIIDHGPEAARGIAVSVDVRYQSKEFAELTCSIMANGIKSIYIK 123
 NTY + KA++ A +++ G A+G+ ++YD R++S EFA + +GIK+Y+++
 Sbjct: 61 NTYTIIRKSEGFARYLLBQGEHVKAGGVVIAYDSRHKSPFAREAAITIGHGKIKAYLFE 120

25 Query: 124 GIRPTMCSYAIRALGCVSGVMITASHNPQAYNGYKAYKEGSGQLDDIADQIANHMDAI 183
 +RPT S+A+R LG G++ITASHNP YNG+K Y +G Q+ +A++ ++ I
 Sbjct: 121 ELRPTPELSFAVRKLGAGGIVITASHNPPEYNGFYQSGDQQLPEPAPNLVKFVNI 180

30 Query: 184 TDYQQIKQIPFEALAGSGSASYIDESIEBAYKGEVLGLTINDTHID--KSVRVVYVTLN 240
 D I E +G+ I E ++ AY + + +N ++ K VR+V+TPL+
 Sbjct: 181 EDELVI PVGERELKGNOTLEMIGEEVDVATHEALKTIIINPELEAGAKDVRIVFTPLH 240

35 Query: 241 GVGNLVPRVLRRRGFENVVYVPEQEMPDQFTTVGVNPEVFKAFAYSESIGKSVDAI 300
 G NLPVR VL GFENV VV EQE+PDP F+TV NPPE AFA + GK +A+
 Sbjct: 241 GTANLPRVRVLEAVGFENVTVKQELPDQFSTVKA/NPEHEAALAEYGGKITEADV 300

40 Query: 301 LLATDPDCDRVALEVKDSKEVIFLNGNKIGALLSYVIPSQRCAICNLPHHPVLVKEIVT 360
 L+ATDPD DRV + V++ GEYI L GN+ G L+ +Y+ SQ+ G LP + + +K+IVT
 Sbjct: 301 LIATDPDADRGVAVQNGAGEVIVLTGNQTGGLMLHYLLSQKKEGQLPVGIALKTIIVT 360

45 Query: 361 GDLSKVIADKNIETVETLTGFKNIOGKANEYDISDKTYLPGYEESIGFCYGTFFVRDKD 420
 + + LA+ + I V+TLTGFK I K EY+ S+ +LPGYEES G+ G FVRDKD
 Sbjct: 361 SEFGRALAEFDGIPMDVTLTGFKF IGKIKIKEYBQSGEHQFLPGYEESYGLIGDFVRDKD 420

50 Query: 421 AVSASMGVREMTAYYKERGCTLLDLVQTIYDKPGYMYERQPSLELGAEGQERISIMED 480
 AV A ++ EMTAYYK RG TL D L ++D++GYE E S+ L+G G E+I ++
 Sbjct: 421 AVQACLLAAEETAYYKSGMTLVDGLLELDFRQYGYRGLTSTILKSGVGEKIQIVLQ 480

55 Query: 481 FRQDPILQVGEHTLENSIDFKDGYK-----DFFQKNCIAKYFNEGSGNYALRPSG 529
 FRQ P QV + + D++ K P N LKY +GSW+ LRPSG
 Sbjct: 481 FRQSPPKQVNDQQVVVIEDYQTKKESVVKERTVEALTLPTNKLKYMLEDGSGNFCALRPSG 540

Query: 530 TEPKIKCY 537

TEPK+K Y

50 Sbjct: 541 TEPKLIK Y 548

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 265> which encodes the amino acid sequence <SEQ ID 266>. Analysis of this protein sequence reveals the following:

Possible site: 35

55

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.5497 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

60

An alignment of the GAS and GBS proteins is shown below:

Identities = 470/564 (83%), Positives = 517/564 (91%)

Query: 1 MSHMYKRIYQEWLENDLSLGDIKSDLSAIGDESEIQDRFYKTLERGTAGLRLGAGT 60
 MS+M Y E+YQEWL N+ L DIK+DL A+K +E+SIQDRFYKTLERGTAGLRLGAGT
 5 Sbjct: 1 MSNMTYNEVYQEWLHNNDLSDDIKADLAIKDNEAEIQDRFYKTLERGTAGLRLGAGT 60

Query: 61 NRMNTYVGVKAAQALANTIIDHGPEALARGIYVSYDVRYSKFEALTCISIMANGIKSY 120
 NRMNTYVGVKAAQALANTIIDHGPEA+ +GIYVSYDVRYS+ FAELTCISIMANGIK+Y
 10 Sbjct: 61 NRMNTYVGVKAAQALANTIIDHGPEAVKKGIAVSYDVRYSRTEFAELTCISIMANGIKAY 120

Query: 121 IYKGIPTPMCSYAIRALGCVSGVMITASHNPQAYNGYKAYKBEQSQILDDIADQIANHM 180
 +YKGIPTPMCSYAIRALGCVSGVMITASHNPQAYNGYKAYW+BQSQILDDIADQIA HM
 Sbjct: 121 IYKGIPTPMCSYAIRALGCVSGVMITASHNPQAYNGYKAYWQBGSQILDDIADQIAQHM 180

Query: 181 DAITYQOIKQIPFEALASGSASTIDESIEBAYKKEVLGLTINDTIDKSVRVVYTPLN 240
 A+T YQ+IKQ+PFE+AL SG +YIDESIEBAYKKEVLGLTINDTIDKSVRVVYTPLN
 15 Sbjct: 181 AALTQYQIKQMPFEALDSGLVITYIDESIEBAYKKEVLGLTINDTIDKSVRVVYTPLN 240

Query: 241 GVGNLPRVRLRRRGFENVVYVPEQRMPPDPFTTVGYNPEVPKAFAYSESLGKSVADAI 300
 GVGNLPRVRLRRRGFENVVYVPEQRMPPDPFTTVGYNPEVPK FAYSE LK+VADAI
 20 Sbjct: 241 GVGNLPRVRLRRRGFENVVYVPEQRMPPDPFTTVGYNPEVPKTFAYSEKLKAVADAI 300

Query: 301 LLATDPDCDRVALEVKDSKGEYIFLNGNKIGALLSYIIFSCRCALGNLPHFVLVKSIVT 360
 L+ATDPDCDRVALEVK++ G+Y+FLNGNKIGALLSYIIFSCRCALGNLPHFVLVKSIVT
 25 Sbjct: 301 LLATDPDCDRVALEVKHAGVYVFLNGNKIGALLSYIIFSCRCALGNLPHFVLVKSIVT 360

Query: 361 GDLKSVIADKYNITVETITGFIGNIGKANEYDLSKDKTYLFGYEESIGFCYGTVPVRDKD 420
 GDLK+ IA Y IETVETITGFIGNIGKANEYD++K K YLFGYEESIGFCYGTVPVRDKD
 Sbjct: 361 GDLGRAIASHYGIETVETITGFIGNIGKANEYDVTKQKNLFGYEESIGFCYGTVPVRDKD 420

Query: 421 AVSASWVVMETAYYKRGQTLLDVLQTIYDKFGYYNERQPSLELGAGQERISIMED 480
 AVSASW+VEM AYYK++GQ LLDVLQITY FGYYNERQ +LELEG EGQ+RI+RIMED
 30 Sbjct: 421 AVSASWIVMEMAAYYKKQNLDDVLQTIYATFGYYNERQIALELEGIGQRIARIMED 480

Query: 481 FRQDPILQVGMTLENSIDFKGVKDPFKCNCLKYVFBGSWYALRPSGTEPKIKCYLYT 540
 FRQ PI V RM L+ +IDF DGY+DFPKCNCLK+Y +GSWYALRPSGTEPKIK YLYT
 35 Sbjct: 481 FRQTPIASVAERMLDKTIDFIDGYQDFPKCNCLFYLLDGSWYALRPSGTEPKIKFYLYT 540

Query: 541 IGCTEADSLSKINATESACRAKGN 564
 IG T+ +S +KL+ATE+ACR K+N
 40 Sbjct: 541 IGQTGRNSATKLDATAEACRTKIN 564

Based on this analysis, it was predicted that these proteins could be useful antigens for vaccines or diagnostics.

45 Example 80

A DNA sequence (GBSx0080) was identified in *S. galactiae* <SEQ ID 267> which encodes the amino acid sequence <SEQ ID 268>. This protein is predicted to be methylenetetrahydrofolate dehydrogenase (fold). Analysis of this protein sequence reveals the following:

Possible site: 48
 50 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.4672 (Affirmative) < succ>
 55 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

60 >GP:AA044612 GB:U58210 tetrahydrofolate dehydrogenase/cyclohydrolase
 [Streptococcus thermophilus]
 Identities = 209/282 (74%), Positives = 248/282 (87%)

Query: 1 MTELIDGKALSQMKAELGRKVERLKBQHGIIPLGLAVILVGDNPASQVYVRNKERSALEA 60
 M ++DGKAL+ MQ +L KV RLKE+ I+PGL VI+VG+NPASQVYVRNKER+ A +A
 Sbjet: 1 MAITMDGKALAVNMQEQQLQERVARLKEKEWIVPGLVVMVGENPASQVYVRNKERAKKA 60

Query: 61 GFKSETRLRSEISIQESELIDIIHOYNEDKSIHGILVOLPLPQHINDKKI ILAIDPKKVD 120
 GF +T+ LSEIS+EEELI+I +YN++ HGILVOLPLP HIN+ +I+LAIDPKKVD
 Sbjet: 61 GFKSKTVNLSEISIEEELIEVIEKYNQNPPLPHGILVOLPLPNHINBMRILLAIIDPKKVD 120

Query: 121 GFHPMTNTHLMSGRPMVPCPAGIMEMFREYHVDLEGKHAVIIGRSNIVGKPMQALLLD 180
 GFHPMTG+LN+GRP MVPCPAGIME+ REY+V+LEGG AVTIGRSNIVGKPMQALLL+
 Sbjet: 121 GFHPMTGNLNGRPQMVPCPAGIMEILREYNVELEGKTAVTIGRSNIVGKPMQALLLE 180

Query: 181 KNATVTLTHSRFTNLSEVTKRADILIVAIGQGHFVTKDFVKEGAVVIDVGNRRDENGKLI 240
 KNATVTLTHSFT +L+V +AD+LIVAIG+ FVT++FVKEGAVVIDVG+NRDE GKL
 Sbjet: 181 KNATVTLTHSRTPHLAKVQNKADVLIIVAIGRAKFTERFVKEGAVVIDVGINRDEBCKLC 240

Query: 241 GDVVFQVQAEVASMITPVGQVGPMTITMLLEQTYQAAALRSV 282
 GDV F+QV E SMITPVGQVGPMTITML+RQTYQAAALRS+
 Sbjet: 241 GDVDFDQVKEKVSMTITPVGQVGPMTITMLBQTYQAAALRS 282

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 269> which encodes the amino acid sequence <SEQ ID 270>. Analysis of this protein sequence reveals the following:

Possible site: 22

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3368(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 230/281 (81%), Positives = 257/281 (90%)

Query: 1 MTELIDGKALSQMKAELGRKVERLKBQHGIIPLGLAVILVGDNPASQVYVRNKERSALEA 60
 MTELIDGKAL+QMKQ RL KV LK++ GI+PGLAVILVGD+PASQVYVRNKER+AL
 Sbjet: 3 MTELIDGKALAQMKQELAAKVNRLKQKKGIVPGLAVILVGDNPASQVYVRNKERAAALTV 62

Query: 61 GFKSETRLRSEISIQESELIDIIHOYNEDKSIHGILVOLPLPQHINDKKI ILAIDPKKVD 120
 GFKSET+RLSE I QRELI +I +Y N D +THGILVOLPLP HINDKKI ILAIDPKKVD
 Sbjet: 63 GFKSETVRLSEFIQRELIHAVIERYNADNTIHGILVOLPLPQHINDKKI ILAIDPKKVD 122

Query: 121 GFHPMTNTHLMSGRPMVPCPAGIMEMFREYHVDLEGKHAVIIGRSNIVGKPMQALLLD 180
 GFHPMTNTHLMSGRP+MVPCP+GIME+ REY+V+LEGGHAVIIGRSNIVGKPMQALLLD
 Sbjet: 123 GFHPMTNTHLMSGRPLMVPCP+PSGIMELREYNVNLEGGHAVIIGRSNIVGKPMQALLLD 182

Query: 181 KNATVTLTHSRFTNLSEVTKRADILIVAIGQGHFVTKDFVKEGAVVIDVGNRRDENGKLI 240
 KNATVTLTHSRTR L EV + AD+LIVAIGQGHF+TK ++K+GA+VIDVGNRRD+NGKLI
 Sbjet: 183 KNATVTLTHSRTRQLEVEVRCADVLIIVAIGQGHFTKQYIKDGAVIDVGNRRDENGKLI 242

Query: 241 GDVVFQVQAEVASMITPVGQVGPMTITMLLEQTYQAAALRSV 282
 GDV F++VARVA+ ITPVGQVGPMTI MLLEQTYQAAALRS
 Sbjet: 243 GDVAFDEVASVAAKITPVGQVGPMTIAMLLEQTYQAAALRS 283

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 81

A DNA sequence (GBSx0081) was identified in *S.galactiae* <SEQ ID 271> which encodes the amino acid sequence <SEQ ID 272>. Analysis of this protein sequence reveals the following:

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Possible site: 39

>>> Seems to have no N-terminal signal sequence

INTEGRAL Likelihood = -3.24 Transmembrane 39 - 55 (38 - 58)

----- Final Results -----

bacterial membrane --- Certainty=0.2296(Affirmative) < succ>

bacterial outside --- Certainty=0.0000(Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 9623> which encodes amino acid sequence <SEQ ID 9624> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:AA044613 GB:U58210 orf1091 [Streptococcus thermophilus]

Identities = 149/277 (53%), Positives = 191/277 (68%)

Query: 1 MIVGQBARALIKPRPKSSHKGDIYGVLLIGFFYPYGGAIINAAACVKGAGLVTVATQ 60

M V + R + I + PR + SHEG YG VLL + GG YFYGGAIINAA + ACV + GAGLVTVAT

Sbjct: 1 MKVDDDLVRQVIRPRELGSSEKSGYGRVLLVGLLYPYGGAIINAAACVNGAGLVTVATD 60

Query: 61 SCHPIPSHLSQLEPMVAFDSDDYKWLKSIQVSDVIVIGPGLGVSESRKILNQTMEKIQS 120

NI + LH + LPE MAFD + + + + DVI + IG GLG E + + L + I + S

Sbjct: 61 RENIIALHAHLPEMAFDLRETERFLDKLEAADVILIGSLGSEETADWALELVLNARS 120

Query: 121 HQSVILDGSAITLSSGAFQTKAGNLVLTTPHOKWERLSSGIASVQQTENTOTALKSFF 180

+ Q + + + + DGSAL LL + + + + L + LTTPHOKWERLSS + A + S + Q + NTQ AL + F

Sbjct: 121 NQNLVVDGSAINLLAKNQSSLPKCHLLTTPHOKWERLSSGLAISQSVNTQRALEEFQ 180

Query: 181 KGTILVAKSHSTRIFQDLDEKEIIVGGPYQATGGMDTLCSMIAGMLAQFEASPLDKVS 240

GTILVAKS T + + Q + + VGGPYQATGGVGDITL GN + AG LAQF V

Sbjct: 181 GSTILVAKSHKTAIVYQGAETVILEVGGPYQATGGMDTLAGNVAGFLAQFASTDSYKAVI 240

Query: 241 VGVYILHSAIAQGLSKAYVVLPTTISDEIPKEMARIS 277

V + LHSAIA + + + AYVVLPT IS IP M + LS

Sbjct: 241 VATWLHSAIADNIAENAYVVLPTIRISKATPSWMKKLS 277

No corresponding DNA sequence was identified in *S.pyogenes*.

SEQ ID 272 (GBS413) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 79 (lane 2; MW 34.2kDa). It was also expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 171 (lane 7; MW 59kDa).

GBS413-GST was purified as shown in Figure 218, lane 12.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 82

A DNA sequence (GBSx0082) was identified in *S.agalactiae* <SEQ ID 273> which encodes the amino acid sequence <SEQ ID 274>. This protein is predicted to be Exonuclease VII large subunit (xseA). Analysis of this protein sequence reveals the following:

Possible site: 36

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3172(Affirmative) < succ>

bacterial membrane --- Certainty=0.0000(Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

5 >GP:CA14361 GB:Z99116 similar to exodeoxyribonuclease VII (large
subunit) [Bacillus subtilis]
Identities = 193/446 (43%), Positives = 283/446 (63%), Gaps = 10/446 (2%)

10 Query: 4 YLSVS/TLTKYLKLPKDPYLERVYL/GQVSNFR-RRPNHQYPSLKDDKSVIQAQWMSH 62
Y++VS LTKY+K KPD DP+LE ++ G+SN + H YP+LK+ K +Q+ M++
Sbjct: 6 YVTVSALTKYIKRGFDVDFELENWIKGELSNVKIHTGRHYFTLKERRKRRKMSVHFARQ 65

15 Query: 63 FKLLGFELEEMKVNWVGRVQVLYEPSPGYSIIVKAEKPDGIGALAIQFBLKKKLQAGY 122
++L F+ E GMKV V G + +YEPSPG+Y + + +PDG+GAL + +E+LKKKL+ G
Sbjct: 66 SERLPFKFENGKMKVLRGGISVYEPSPGYSIIVKAEKPDGIGALAIQFBLKKKLQAGY 125

20 Query: 123 FDDRHKQLIPQFVRKIGVVISPSGAVIRDIITVSRFPFGVEILLFPTKVGQEGAAQEI 182
FDDR+K+ IP F IGWVSP+GA +RD+ITT+ RR+P V++++ P VQGE A++ I
Sbjct: 126 FDRYKQKQIPAFATIGVVISPTGA+VRVITTLKRRYPLVKYIVLPAWQEGASRSIV 185

25 Query: 183 OTIALANEKIDLLIVGRGGGSIEDLWAFNEECVVEALPESRLPVISSVGHSTDTTLD 242
I ANEK+ D+LIVGRGGGSI+LWAFNEE V AIF S +P+Is+VGHST T+D+D
Sbjct: 186 TRIEANEKEICDVLIVGRGGGSIEDLWAFNEEIVARAIFASNPIIISVGHSTDTTID 245

30 Query: 243 FVADRRAAATPTAAAEATPVTKIDILSNWITRENNMYQSSLRILIRTKERLQSKQSVIF 302
FVAD RAATPT AAE+A P T D++ E RM ++ + + ++ R+Q + S F
Sbjct: 246 FVADIRAATPTGAABIAVPH-TDLIERKTAEVMTAMQCHQLQGEKRIQTLQSSYAF 304

35 Query: 303 RQPERLYDGFQKLD---NLNQOLTYSMRDKLQTVRQKQGLLHQKLGIDLQRIHIYQ 358
R P+RLY Q+ D QLT + K + + + L LKQ YQ
Sbjct: 305 RPFKRLYQKQKQDFLAYQQFQAQTLALDERSKQLERETRYLEALHPHEQLQDARTRY 364

40 Query: 359 ERVQSRRLSSTMTSYDSEKLARPEKAQDALISDSRIVARGYALIEKHNTLVSTING 418
E+ Q R+ M Q ++F+ L +L +++ ROY++ K L+ + +
Sbjct: 365 EQTNQLRK---NNHQMQLHSQFQTVLGLKMLASPLQVMERGYSLAYEKDKLKSVSQ 420

45 Query: 419 INEGDHQVMDOLLEVEVKVROE 444
I E D L++K++DG+L EV + R E
Sbjct: 421 IREQRLEIKLDGVLTCVLEKRG 446

40 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 275> which encodes the amino acid
sequence <SEQ ID 276>. Analysis of this protein sequence reveals the following:

Possible site: 61

>>> Seems to have no N-terminal signal sequence

45 ----- Final Results -----
bacterial cytoplasm --- Certainty=0.3275 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

50 An alignment of the GAS and GBS proteins is shown below:

Identities = 321/446 (71%), Positives = 386/446 (85%)

55 Query: 1 MSDVLSVSTLTKYLKLPKDPYLERVYL/GQVSNFRRRPNHQYPSLKDDKSVIQAQWMS 60
M+DYL+V+ LTKYLKLPD+DPYLERVYL/GQVSNFR+RP HCYFSLKD+ +VIQAQW+
Sbjct: 6 MDVLTVTHTLTKYLKLPKDPYLERVYL/GQVSNFRKRPKPTHYPSLKDDKSVIQAQWMS 65

60 Query: 61 GHFKKLGFELSEGMKVNWVGRVQVLYEPSPGYSIIVKAEKPDGIGALAIQFBLKKKLQ 120
G +KKLGF+LESGMK+NV+GRVQVLYEPSPGYSI+ +EKAEKPDGIGALA+QFBLKKKL+
Sbjct: 66 GYVKKLGFDELSEGMKINWVGRVQVLYEPSPGYSIIVKAEKPDGIGALAIQFBLKKKLQ 125

65 Query: 121 GYFDRHKQLIPQFVRKIGVVISPSGAVIRDIITVSRFPFGVEILLFPTKVGQEGAAQ 180
GYP+ +HKQ +POFV KIGV+TSPSGAVIRDIITVSRFPFGVEILLFPTKVGQ+GAAQ
Sbjct: 126 GYFDRHKQLIPQFVRKIGVVISPSGAVIRDIITVSRFPFGVEILLFPTKVGQEGAAQ 185

```

Query: 181 IAQTALANEEKKDLILLVGRGGGSIEDIMAFNEECVVEAIFESRLPVISSVGHETTTL 240
      + I AN+++DLDMIVGRGGGSIEDIMAFNEE VV+AIRES+LPVISSVGHETTTL
Sbjct: 186 VVANIRRRANQREDLILLVGRGGGSIEDIMAFNEEIVVQAIRESQLPVISSVGHETTTL 245

Query: 241 ADFVADRRRAATPTAAAEIATPTIKIDILSWITERENRMVQSSRLRLIRTKERLQKSKQSV 300
      ADFVADRRRAATPTAAAEIATPTIK D+SWI HR+NR YQ+ LR I+ ++E + K QSV
Sbjct: 246 ADFVADRRRAATPTAAAEIATPTITKIDLMSWITERVQNRSYQACLRRIKQEQWVDKLSQSV 305

Query: 301 IFRQFERLYDGFLOKLENLNQCLTYSMRDKLQTVRQKQELLEQKQKIDLQRIHIYQER 360
      IFRQFERLYD +IQ+D L+ L +M+D+L + ++ + L L L+ +I YQ+R
Sbjct: 306 IFRQFERLYDAYLQKIDRLSMITLNTNMKDRISGAKENQVLQHALANSQ/LQTKIERYQDR 365

Query: 361 VVQSRRLSSNTTSQYDSKLARFEKAQDALISLDSRIVARGYALIEKNHILVSTTNGIN 420
      V ++RL + M SQYDS+LARFEKAQDAL+SLD+SRI+ARGYA+IEKN LV++ + I
Sbjct: 366 VATAKRLLMANNSQYDSQLARFEKAQDALISLDSRIIARGYAMIEKNQALVASVSQIT 425

Query: 421 BGDHLQVNMQDGLLEVEVQDVRCENI 446
      +GD L +RM+DG L+VEVQDV+ ENI
Sbjct: 426 KGDQLTINMRDQGLDVEVQDVRCENI 451

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 83

- 25 A DNA sequence (GBSx0083) was identified in *S.agalactiae* <SEQ ID 277> which encodes the amino acid sequence <SEQ ID 278>. Analysis of this protein sequence reveals the following:

Possible site: 33

```

>>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.2913 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP: AAG07429 GB:AE004821 exodeoxyribonuclease VII small subunit
[Pseudomonas aeruginosa]
Identities = 26/66 (39%), Positives = 51/66 (76%), Gaps = 2/66 (3%)

Query: 1 MSDKKT--FEENLQELTIVSRLLTGDEVALEDAIEFQKGMILISKELQRTLKRAETLVK 58
      M+ KKT FE++L EL+T+V RLK+G+++LE++ F+G+ +++E Q +L +AE+ +
Sbjct: 1 NARKKTLDFEQSLATLQTLVERLESSELSESLGAFEGGIRLTRECQTSLSQAQKVAI 60

Query: 59 VMQADG 64
      +++ DG
Sbjct: 61 LLERDG 66

```

- 50 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 279> which encodes the amino acid sequence <SEQ ID 280>. Analysis of this protein sequence reveals the following:

Possible site: 51

```

>>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.2796 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

- 60 An alignment of the GAS and GBS proteins is shown below:

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Identities = 55/70 (78%), Positives = 65/70 (92%)

Query: 1 MSKDKTFEENLOELETIVSRLETDGVALEDATAEPQKQMLISKELQRTKKARETLVQVM 60
 MS KTFEENLQ+LETTIV+LE GDV LE+AI+EPQKQML+SKELQ+TL+AE+TLVQVM
 Sbjct: 1 MSKDKTFEENLOELETIVSRLETDGVALEDATAEPQKQMLISKELQRTKKARETLVQVM 60

Query: 61 QADGTEVEMD 70
 QADGTEV+MD
 Sbjct: 61 QADGTEVEMD 70

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 84

A DNA sequence (GBSx0084) was identified in *S.galactiae* <SEQ ID 281> which encodes the amino acid sequence <SEQ ID 282>. Analysis of this protein sequence reveals the following:

Possible site: 58

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2614(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:BAA25265 GB:AB003187 farnesyl diphosphate synthase [Micrococcus
 luteus]
 Identities = 126/258 (48%), Positives = 175/258 (66%), Gaps = 2/258 (0%)

Query: 27 LKALYVSVDGGKRRIRPRLLEILLEGFOVELIDGHYDVAAALEMINTGSLIHDDLPA 86
 L +AI YS+ GKKRIRP +L L+ G DG ALEMINT SLIHDDLPA
 Sbjct: 31 LREAINYSLSAAGKRRIRPRLVLTLDLGGNARDG-LPFGIALEMINTYSLIHDDLPA 89

Query: 87 NDDPRGRRLTNHKKFDEATAVLAGDSLELDPPDLVKGAGFKADVTVRLISLMSAGSFG 146
 NDD+RRG+LTNHK+FDEATA+LAGD+L D F ++ A++ + LI LLS +GS G
 Sbjct: 90 NDDYRRGKLTNHKKFDEATAVLAGDALLTDAFQCLINTQINAKIKSLINLLSTASGNG 149

Query: 147 MVGGQMLDMKGENKVLSDLSLHINKTGRLAYPPVAAAGILAKSEEVKGLHQA 206
 MV GQMLDM+GE+K L++++L IHI+KTG L+ V+AGI+ ++ +L+ G
 Sbjct: 150 MVYGGQMLDMQGHKLTITNLERIHIHKTGELIRAAIVSAGIIMNNDAGIKQLNIIGN 209

Query: 207 IGHAFOVRDILDLVTASPERLAKTPNKKIVAKCTTYRNLLGLDKSQRLIDTLKQQAIF 266
 +G FQ++DDILDV SFE +GKT D+ +K+TY +LLGL+ S+++L+D L +
 Sbjct: 210 VGLMFQIKDDILDVSGSPENIGKTVSGSLNNDKSTVSVSLGLEASQLLNDKLTITYDAL 269

Query: 267 QNLKKNFANKRIIDII 284
 + L+ N N + + I I
 Sbjct: 270 KTLQ-PINDNIAKTLITYI 286

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 283> which encodes the amino acid sequence <SEQ ID 284>. Analysis of this protein sequence reveals the following:

Possible site: 38

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3887(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 192/289 (66%), Positives = 237/289 (81%)

```

5 Query: 2 MVTIKKIDEAIIHRYKQTHSVSPDLIKALLYSVDGSGKRIRPRILLILBEGFVGLIDG 61
      M + +IDEAI RYK T + VS +LI AILYSVD GGRIRP IILE++BEGFV L +
      Sbjct: 1 MDKILARIDEAIRRYKTTNSGVSEELDAILYSVDGSGKRIRPILILEMIEBEGFVSLQNA 60

10 Query: 62 HYDVAALAEIMHTIGSLIHEDLEAMUNDGFRGRITNHHKPFDEATAVLACDLSFLDPFDLV 121
      R+DAAALAEIMHTIGSLIHEDLEAMUNDG+RGRITNHH+ P EATA+LACDLSFLDPF L+
      Sbjct: 61 HFDLAALAEIMHTIGSLIHEDLEAMUNDGFRGRITNHHKQGEATAVLACDLSFLDPFGLI 120

15 Query: 122 VKAGFKADVTVRLIELLSMSGSGSPGVNGGQMLDMKGENKVLISIDDSLIHINKTGRLAY 181
      +A + +V V LI+ LS++G+PGNVGGQMLDMKGEN+ LS+ LSLIH+NKTG+LLA+
      Sbjct: 121 AQAEINSEVKVALIQELSLASTPGNVGGQMLDMKGENQALSLPOLSLIHINKTGRLAF 180

20 Query: 182 PFVAAGILAEKSEEVKGLHOGGLIGHAFQVRDDILDVTASFEELGKTPNDIVAETK 241
      PF AA ++ E++ V+ +L QAG+LIGHAFQ+RDDLVDVTASPE+LGKTP ND+ AEK T
      Sbjct: 181 PFKAALITHEQMTVRQQLSQAGMLIGHAFQIRDDILDVTASFEDLGKTPKDLFAEKAT 240

20 Query: 242 YPNLLGLDKSQEILDOTLKQAIFONLEKKKAMFNARKIID:IEGLRLN 290
      YP+LLGL+ S ++L ++L +A IPQ LE F + I +IEGLRLN
      Sbjct: 241 YPSLLGLEASTQLLTSLDQALITFQTLSEVGVFPQIITKGLIEGLRLN 289
  
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 85

A DNA sequence (GBSx0085) was identified in *S. galactiae* <SEQ ID 285> which encodes the amino acid sequence <SEQ ID 286>. This protein is predicted to be hemolysin-like protein (tly). Analysis of this protein sequence reveals the following:

```

30 Possible site: 37

    >>> Seems to have no N-terminal signal sequence
    INTEGRAL    Likelihood = -0.75    Transmembrane    152 - 168 ( 151 - 168)

35 ----- Final Results -----
      bacterial membrane --- Certainty=0.1298 (Affirmative) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

40 The protein has homology with the following sequences in the GENPEPT database:

    >GP:BAB06497 GB:AP001516 hemolysin-like protein [Bacillus halodurans]
    Identities = 162/270 (60%), Positives = 202/270 (74%), Gaps = 3/270 (1%)

45 Query: 3 KERVDVLAYKQGLFDTRQAKGGVWAGMVINIGERYDKPGEKVADDTLKLKGEKLY 62
      KERVDVL ++GL +TRB+AKR +MAG+V + ER DKPV KV DT L +XGE L Y
      Sbjct: 4 KERVDVLLVERGLMETREKAKRSTIMAGLVFS--GHERVDKPKLVDRDTPLSVGEVLFPY 61

50 Query: 63 VSRGGLKLEKALQVFEISVADKLTIDIGASTGGFTDVMQLSGARLVYAVDVGTNTQVWL 122
      VSRGGLKLEKAA+ F++ + D++ +DIGASTGGFTD LQ+GA VYAVDVG NQL WKL
      Sbjct: 62 VSRGGLKLEKATRAFDLHLDTRVVLIDIGASTGGFTDCALQNGATFYAVDVGVYGNLAWKL 121

55 Query: 123 RQDRHVRSMBCQYNFRYAQKEDPKKGLPEFASIDVSFISINLLPALKEILVDGSGQVALI 182
      RQD RV ME+ NFRY + E + GLP A+IDVSFISL LILP LK +L++ VVAL+
      Sbjct: 122 RQDERVVVMKRNFRYIKPEVLKGLPMATIDVSFISLKLILPVLLKMLGNSDVVALV 181

60 Query: 183 KPQFEAGREIQIGKIGVNDKLVRHKKVLTITVPNTKDYGVTVKHLDFSPIQGKHGNIPLM 242
      KPQFEAGRE++GK GIV+DK VR+KVL+T+ P GY V LDFSPI GG GNIEPL+
      Sbjct: 182 KPQFEAGREEVGKKGIVRDKSVHKKVLTIVEFALKRGYAVGGLDFSPITGSGENIPLL 241

60 Query: 243 HLQKQDDPQNLV-LDQIQDVIEKAHKRFKK 271
  
```

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HL +D ++ + I+D +E+AE E KK
 Sbjct: 242 HLMWRKDKESFISQEMIRDTVERAHLK 271

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 287> which encodes the amino acid
 5 sequence <SEQ ID 288>. Analysis of this protein sequence reveals the following:

Possible site: 37

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -2.92 Transmembrane 150 - 166 (149 - 168)

----- Final Results -----

bacterial membrane --- Certainty=0.2168 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP:BAB06497 GB:AP001516 hemolysin-like protein [Bacillus halodurans]
 Identities = 156/270 (57%), Positives = 196/270 (71%), Gaps = 3/270 (1%)

Query: 3 KERVDVLAYKQGLFETREQAKRGVMAGLVVSVINGQRYDKPGDKIDDGTCLKKGEKLY 62
 KERVDVL ++GL ETR+AKR +MAGLV S +R DKPG K+D T L +KGE L Y
 Sbjct: 4 KERVDVLVERGLMETREKAKRSINAGLVFS--GHERVDKPKGLKVDRTPLSVKGEVLFP 61

Query: 63 VSRGGLKLEKGLHVFVGSVANQIGIDIGASTGGFTDVMQLQDGAFLVYADVGTNQLVWKL 122
 VSRGGLKLEK + F + + + +DIGASTGGFTD LQ+GA VYADVQ NQL WKL
 Sbjct: 62 VSRGGLKLEKALRAFDLHLTRVDVLDIGASTGGFTDCLQNGATFVYADVGVNQLAKWL 121

Query: 123 RODPRVRMBQYNFRYAQPEDFNBOQVPFASIDVSPISLSLILPALHNVLSDQGVIALI 182
 ROD RV ME+ NFRY +PE G P A-IDVSPISL LIIP L +L + V+AL+
 Sbjct: 122 RODPRVVMERTINFRYLKPEVLEKGLPMATIDVSPISLKLILPLKMLLENSDVVALV 181

Query: 183 KPQFEAGREQIGKKGIVKDKQIHEKVIQKVMDFASGVGPTVKGLDPSPIQGGHGNIEFLA 242
 KPQFEAGRE++GKKGIV+DK +H+KV+ +++FA G+ V GLDFSPI GS GNIEFL
 Sbjct: 182 KPQFEAGREEVGKKGIVRDKSVKQVLSTIVEFALKEGYAVGGLDPSPIQGGHGNIEFL 241

Query: 243 HLAKSQTPET-LAPHLIQKVAKAHKEFEK 271
 HL + E+ ++ +I+ V +AH E +K
 Sbjct: 242 HLMWRKDKESFISQEMIRDTVERAHLK 271

An alignment of the GAS and GBS proteins is shown below:

Identities = 214/275 (77%), Positives = 238/275 (85%)

Query: 1 MAKERVDVLAYKQGLFETREQAKRGVMAGLVVSVINGQRYDKPGKGVADDTCLKKGEK 60
 M KERVDVLAYKQGLF+TRQAKRGVMAG+V++VING+RYDKPG+K+ D TELKLGKGL
 45 Sbjct: 1 MKERVDVLAYKQGLFETREQAKRGVMAGLVVSVINGQRYDKGDKIDDGTCLKKGEK 60

Query: 61 KYVSRGGLKLEKALQVFEISVADKLTIDIGASTGGFTDVMQLQSGARLVYADVGTNQLVW 120
 KYVSRGGLKLEK L VF +SVA++ IDIGASTGGFTDVMQLQ GA+LVYADVGTNQLVW
 Sbjct: 61 KYVSRGGLKLEKGLHVFVGSVANQIGIDIGASTGGFTDVMQLQDGAFLVYADVGTNQLVW 120

Query: 121 KLRQDRVRMBQYNFRYAQKEDFKGLPEFASIDVSPISLSLILPALKEILVGGGVQA 180
 KLRQ DRVRMBQYNFRYAQ EDF EG P FASIDVSPISL+LILPAL +L D GQV+A
 Sbjct: 121 KLRQDRVRMBQYNFRYAQKEDFNBOQVPFASIDVSPISLSLILPALHNVLSDQGVIA 180

Query: 181 LKIPQFEAGREQIGKKGIVKDKLHVRKVLTVINFTKDYQYTVKHLDSPSQGGHGNIEF 240
 LKIPQFEAGREQIGK GIVKDK +HKV+ V +F YG+TVK LDSPSQGGHGNIEF
 Sbjct: 181 LKIPQFEAGREQIGKKGIVKDKQIHEKVIQKVMDFASGVGPTVKGLDPSPIQGGHGNIEF 240

Query: 241 LHLKQKQDPONLVDQIQDVIEKAHKEFKNEE 275
 L HL K Q P L IQ V+ KAHKEF+K+E+E
 Sbjct: 241 LAHLAKSQTPETLAPHLIQKVAKAHKEFEKKEE 275

SEQ ID 286 (GBS310) was expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 57 (lane 3; MW 34kDa). It was also expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 61 (lane 4; MW 58.8kDa).

The GBS310-GST fusion product was purified (Figure 210, lane 10) and used to immunise mice. The resulting antiserum was used for FACS (Figure 282), which confirmed that the protein is immunoreactive on GBS bacteria.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 86

A DNA sequence (GBSx0086) was identified in *S. agalactiae* <SEQ ID 289> which encodes the amino acid sequence <SEQ ID 290>. Analysis of this protein sequence reveals the following:

Possible site: 18

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1966 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAA09426 GB:J010954 arginine repressor [Bacillus
stearothermophilus]

Identities = 49/153 (32%), Positives = 84/153 (54%), Gaps = 4/153 (2%)

Query: 1 MKKSERLNLIKQIVLHNAVETQHELLERLEAYGVTLTQMTISRDNNEIGIKVPSAKGRY 60
M K +R I ++I ++NH +ETQ EL+ L+ G +TQMT+SRD+ E+ ++KVF A GRV
Sbjct: 1 MNGQRHRIKIREIIMHEIETQDELVDMLKAGFNVTQATVSRDIKELQVKKVPMANGRY 60

Query: 61 IYGLSNENDPIFTTAVAKPIKTSILSISDKLLGLEQFININVI PGNQLIKTFIMSHCQE 120
Y L +D F + +K +++ KL G + + +PGN+ I + +
Sbjct: 61 KYSL--PSDGRFNP--TQKLKRALMDAPVKLDGSGNLLVLKTLPGNAHAIGVLLDNLWN 116

Query: 121 HIFSLTADNNSLLLTARSEADADHQRSMIAML 153
I D++ L+I ++ DA+ + ++ML
Sbjct: 117 EIVGTTCGDITCLICRTAEADKVSQQLGLML 149

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 291> which encodes the amino acid sequence <SEQ ID 292>. Analysis of this protein sequence reveals the following:

Possible site: 50

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1717 (Affirmative) < succ>

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 87/154 (56%), Positives = 118/154 (76%), Gaps = 1/154 (0%)

Query: 1 MKKSERLNLIKQIVLHNAVETQHELLERLEAYGVTLTQMTISRDNNEIGIKVPSAKGRY 60
MKKSERL LK+VL H +ETQH+LIR L +G+ LTQMTISRDNNEIGI+K+PS GRV
Sbjct: 12 MKKSERLELIKQIVLHNAVETQHELLERLEAYGVTLTQMTISRDNNEIGIVKIPSGSGRY 71

Query: 61 IYGLSNENDPFTTAVAKPIKTSILSISDKILGLEQFININVIPGNSOLIKTFIMSHCQE 120
 IYGLS ++ + IK++IL++SDEK GLEQ + + V+PGNS+LIK +++++ +
 Sbjet: 72 IYGLSQDSGKKIVQG-PRSIKSTTLAVSDKTKGLEQLYLYKVVFGNSKLIKRYLLADFSK 130

Query: 121 HIFSLTADNSILLIAXSEADADHRCQMIAMLE 154
 IFSL ADD+SLIAXS ++AD IRQ ++ ++
 Sbjet: 131 AIFSLIADDSILLIAXSPSEADMIRQETILLMMQ 164

- 10 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 87

A DNA sequence (GBSx0088) was identified in *S.galactiae* <SEQ ID 293> which encodes the amino acid sequence <SEQ ID 294>. Analysis of this protein sequence reveals the following:

- 15 Possible site: 15
- >>> Seems to have no N-terminal signal sequence
- Final Results ----
- 20 bacterial cytoplasm --- Certainty=0.3339 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

- 25 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 88

A DNA sequence (GBSx0089) was identified in *S.galactiae* <SEQ ID 295> which encodes the amino acid sequence <SEQ ID 296>. This protein is predicted to be DNA repair protein recN (recN). Analysis of this protein sequence reveals the following:

- Possible site: 50
- >>> Seems to have no N-terminal signal sequence
- Final Results ----
- 35 bacterial cytoplasm --- Certainty=0.1651 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
- 40

The protein has homology with the following sequences in the GENPEPT database:

>GP: CAB14355 GB:Z99116 recN [Bacillus subtilis]
 Identities = 244/567 (43%), Positives = 366/567 (64%), Gaps = 18/567 (3%)

45 Query: 1 MLEISIKNFAIIEETISLNFSTGTVLTGCTGAGKSIITIDAMNMLGSRASVEVIRKGN 60
 ML E+SIKNFAIIEE++++FR G+TVLTGCTGAGKSIITIDA++++G R S E +R+G
 Sbjet: 1 MLEISIKNFAIIEETISLNFSTGTVLTGCTGAGKSIITIDNISLVGCRGSEFVRVYGEA 60

Query: 61 KAEIRGFTSVKNGSLVOLLFENGIELADELIIRREIPONGRSVBRINGQMVNLSTLKA 119
 KAE+EG F +E ++ + G EI++DE+I+ RR+I +G+SV R+NG++V +++L+
 50 Sbjet: 61 KAELEGLFLLESHPVLCVCAEQGIDVSDRMVIMRRDISTSGKSVCRVNGKLVITIASLRE 120

Query: 120 VGHVLDIYQCHDQBELMKFNHILMLDFGNTEFNVIKERYQSLFAYQLRIRKRVLDKQ 179
 +G L+DI+QCHD + LM+ H+ +LD+F E + YQ + Y +L K++

-156-

Sbjct: 121 IGRLLLDIHGQHDNQLMEDENHQLQLDQFAGARVESALCTYQSGYQYVKLLKKLKQLS 180

Query: 180 KNEQENKSRIMELFQIARIESVALKSDSDQTLTKQRDKIMNHKNLADTLTNAYMLDNE 239
 ++EQE ++++FC+ RIES L+ +ED+ L +R+ ++ N + I ++L NAY L +E

5 Sbjct: 181 RSBQEMAHCLDLIQFQLEIESAKLNEEDBQLQERQQTINFEKIYESLQAYNALISE 240

Query: 240 EFSSLSNVRSAMNDLMALEEFQREYKDLSTNLSEAYYVIEVTRKLGVDIDDLDFAGLL 299
 + L V A L + + + K +S ++S ++Y++E+ T + + ++D+L+FD L

10 Sbjct: 241 Q-GGLDWGMSAQLEDISDINEPLKMSSEVSNSYLLLEDAITFQMRMLDELEFQPERL 299

Query: 300 QRIENRLDVINTTRKYGGDVNDVLYFDNITKEYSLLTGSESSDALEKELKLEHDLI 359
 IE RL+ I + RKYG V D+L+Y I +E + + +L+KEL + D+

Sbjct: 300 NYIETRLNRIKQLKKYKATVEDILEYASKIIEIDQIENKDSHLQSLKKELOSVGKDA 359

15 Query: 360 ESANQLSLERHKLAKQLENRIQELTLYMEKADPQVQFTKG-----KF 403
 A +S R AK+L +RI +EL LYMKE+ F +F +

Sbjct: 360 VEAANVSQIRKTAAKLADETIRELSLYMEKSTFTDTEFKVITASRNEAPLVNQCPQL 419

Query: 404 NKEGNEIVEFYISTNPGEGFKPLVKVASGGELSRMLAISKAFSRKEKTSIVPDEVDTG 463
 ++G ++V+V+ ISTN GE K L KVASGGELSR+MLAISK FS ++D TSI+PDEVDTG

20 Sbjct: 420 TEQGIDLVKFLISTNPTGEPKLSLKVASGGELSRVMLAISKIFSSQQVTSIIPDEVDTG 479

Query: 464 VSGRVAQALAQKIKIGSGQVLAISHLAQVIALADYQYFIEKISSDSSTVSTVRLLSYE 523
 VSGRVAQALA+KIKH+ QVL I+HL QV A+AD +I K D T +V+ LS +

25 Sbjct: 480 VSGRVAQALAEIKHVSIGSQVLCITLHPQVAAADTHLYIAKELDKGRITRVKLEKQ 539

Query: 524 ERVEEIAKMLAGNNVDTARTQAKELL 550
 E+V EI + +AG VTD + AKELL

30 Sbjct: 540 EKVAEIERSIAGVEVDLTKRHAKELL 566

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 297> which encodes the amino acid sequence <SEQ ID 298>. Analysis of this protein sequence reveals the following:

Possible site: 51

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1215 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 403/550 (73%), Positives = 472/550 (86%)

Query: 1 MLEISIKNPAIIEISLNFETGMVLTIGTGAGKSI IIDAMNMLGSRASREVIRHCAN 60
 MLEISIKNPAII+EISLAF E GMVLTIGTGAGKSI IIDAMNMLG+RAS RVIR CAN

Sbjct: 2 MLEISIKNPAIIEISLNFENGMVLTIGTGAGKSI IIDAMNMLGARASTEVIRHCAN 61

Query: 61 KASIBGFFSVGEKNQSLVQLLEENGIELADHLIRREI PQNGREVSRIKQMNVLSTKAV 120
 KASIBGFFSV+ LV LE +GI + +RLIRK+IF NGRSVSRINGQMNVL+PLK V

50 Sbjct: 62 KASIBGFFVDATPELVACLESSGIAMEEELIRRDIFANGREVSRIKQMNVLATLQV 121

Query: 121 GHYLVDIYQGHQDQELMRPNMHMLDDEFQNTSFNVIKERYQSLFDATYQLRKRVLDKQK 180
 G +LVDI+GQHDQELM+P +H +JD FG+ F +KE YQ +FD Y+ LR++V+DKQK

55 Sbjct: 122 GQFLVDIHGQHDQELMRPQLHQQLDAPDQKAFBQLKENTYQLIPRYKSLRRQVLDKQK 181

Query: 181 NEQENKSRIMELFQIARIESVALKSDSDQTLTKQRDKIMNHKNLADTLTNAYMLDNEE 240
 NE+R+K RI+ML FQIARIE+ AL ED L +RD+LMNHK LADTLTNAY+MLDN+V

60 Sbjct: 182 NEKEHKDRIMLAPQIARIEAALSRGSDORLNQSRDRIMNHQIADTLTNAYVMDNDND 241

Query: 241 PSSLSNVRSAMNDLMALEEFQREYKDLSTNLSEAYYVIEVTRKLGVDIDDLDFAGLL 300
 PSSLSN+RS+MNDL++E+FD EYK +ST+SEAYY++RKY+K+L D ID LDQD G LQ

Sbjct: 242 PSSLSNIRSSNDLSTIQPDSEYKDMSTSI SEAYYILLEVSQKLSDTIDQLDFDQGLQ 301

65 Query: 301 ERIENRLDVINTTRKYGGDVNDVLYFDNITKEYSLLTGSESSDALEKELKLEHDLI 360

EIE RLD++N++TRKYGG+VNDVLDY+DNI KEY LLTG + SS IE ELK LE L+
 Sbjct: 302 EIEFRLDILNSLTKYGGNVNDVLDYDNIKEYQLLTGDDLSGGLEAEKLSIAEKQLVA 361
 5 Query: 361 SANQLSLERHKLAQQLNEIKQELATELYMEKADFQVQFTKGKFNKEGNEIVEFYISTNPG 420
 +A++LS+ RH+LA+QLE EIK EL ELYMEKADF+V FT KFN++GNE +EFYISTNPG
 Sbjct: 362 AASELSVSRHQIARQIARAIKARKLKELYMEKADFVHFFTSKFNRDQNESELYFYISTNPG 421
 Query: 421 EGFKPLVKVASGGGLSRMLIAIKSAFSRKEDKTSIVFDEVDTGVSGRVAQAIAQKIHKIG 480
 EGFKPLVKVASGGGLSRMLIAIK+A SRKEDKTSIVFDEVDTGVSGRVAQAIAQKI+KIG
 10 Sbjct: 422 EGFKPLVKVASGGGLSRMLIAIKAAISRKEDKTSIVFDEVDTGVSGRVAQAIAQKIYKIG 481
 Query: 481 SHQQLVAISHLAQVIATADYQYFIKSSDSSTVSTVRLLSYEEVERIAKRLAGNNVDT 540
 HQQVLAISHL QVIATADYQYFI K S + STVS VRL+ EERVERIA M+AG ++T
 15 Sbjct: 482 RHQQLVAISHLPQVIATADYQYFISKESKEESTVSKVRLLTPEERVERIASTIAGTDMTQ 541
 Query: 541 TARTQAKELL 550
 A TQA+ELL
 Sbjct: 542 AALTQARELL 551

20 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 89

A DNA sequence (GBSx0090) was identified in *S. agalactiae* <SEQ ID 299> which encodes the amino acid
 sequence <SEQ ID 300>. This protein is predicted to be degV protein. Analysis of this protein sequence
 25 reveals the following:

Possible site: 38
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.96 Transmembrane 246 - 262 (246 - 262)
 30 ----- Final Results -----
 bacterial membrane --- Certainty=0.1383 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 35 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:BAB07346 GB:AP001519 unknown conserved protein [Bacillus halodurans]
 Identities = 93/277 (33%), Positives = 152/277 (54%), Gaps = 4/277 (1%)
 40 Query: 1 MSKIKIVTDSGITIEFLIKELDITVVPVLSVMIDGTLSDNDLKAQGFENLMRGSKELP 60
 M+KI IVTDS+ + P+ KRL + VVPLSV+ Y + + +F ++ ++LF
 Sbjct: 1 MTKIAIVTDSITAYLGFPRKAKLGIVTVPLSVFPGFAYQEEVEELSSADFYKELKHIEKLP 60
 45 Query: 61 KTSQPPVGVFARIYEKLMNGBVEHIIAIIHAHTLSTGIE-ASRQGANIAGADVTVIDSTF 119
 TSQP VG+F E +E+L EG E +I+IHL+ +SGT +A G+ + G +V DS
 Sbjct: 61 TTSQPAVGLVFETPERLAKEGGFVVI SIHLKSIIGTQSAITAGSMVGBIEVIGYDSGI 120
 Query: 120 TDQQCKFQVVEAAKLAKEGADLTILARVEEVRQKSELFICGVSTLENLVKGRIGRIVTGL 179
 + + Q V EAAKL KEGAD TI+ ++V+ +++ V L +L +GGR+ +
 50 Sbjct: 121 SCEPQANFVAEAAKLVKEGADPQTII DHLDEVKGTNALPVVHDLSHLHRRGRINAAQV 180
 Query: 180 LSSLINIKVIMELTNHVLPIVVKGR-GLKTFKMLNDNPVBSAQTRKIAETIGSYGKADM 238
 + SLL IK I+ + +VP+ K R K +++ + F E A + + + + D
 55 Sbjct: 181 VGSLLKIKPILHFDGSIIVLEKVRTEKMAARVIGLFAEASSASSVKNTVIHANRLDG 240
 Query: 239 ANNPREKL--AVLGAPISVLETGSIIQTHTGEDAFV 273
 A +++ +B+ G +I TH GE + +
 Sbjct: 241 AEKLADEIRSQPSHVDVSI SHFGPVIGTHLGEISIGL 277

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 301> which encodes the amino acid sequence <SEQ ID 302>. Analysis of this protein sequence reveals the following:

Possible site: 37

```

5  >>> Seems to have no N-terminal signal sequence
    INTEGRAL    Likelihood = -1.54    Transmembrane  180 - 196 ( 180 - 196)
    INTEGRAL    Likelihood = -0.16    Transmembrane  21 - 37 ( 21 - 38)

10  ----- Final Results -----
        bacterial membrane --- Certainty=0.1617(Affirmative) < succ>
        bacterial outside --- Certainty=0.0000(Not Clear) < succ>
        bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

```

15  Identities = 197/279 (70%), Positives = 226/279 (80%), Gaps = 1/279 (0%)

Query: 1  MSKIKIVTDSSTITPELIKELDITVVLPSVMIDGTLVSDNDLKAQGFNLNMRGSKELP 60
      M  IKIVTDSSTITPELIK  LDITVVLPSVMID  LYSNDLK +G FL+LN+ SK LP
20  Sbjct: 5  NGTIKIVTDSSTITPELIKALDITVVLPSVMIDSKLYSDNDLKEBGFHLSLMKASKSLP 64

Query: 61  KTSQPPVGVFARIYEKLMEGVSHI IAIHL/HTLSGTIEASRQAGIAGADVTVIDSTFT 120
      KTSQPPVG+FAE YE L+ +GV I+AIHL+ LSGTIEASRQGA IA A VTV+DS FT
20  Sbjct: 65  KTSQPPVGLFARTYENLVKKGVTDI IAIHLSPALSGTIEASRQAGIAEAPVTVIDSGFT 124

Query: 121  DQCQKFQVVEAAKAKAGADLDITLARVEEVROKSELFVIGVSTLENLVKGGRIGRVTVGL 180
      DQ KFQVVEAAK+AK GA L+ ILA V+ ++ K+EL+ IQVSTLENLVKGGRIGRVTVGL
25  Sbjct: 125  DQAKQFQVVEAAKAKAGASINELIAAQAISKTELYIGVSTLENLVKGGRIGRVTVGL 184

Query: 181  SLLLNKIVMELTNHSLVPIVKGRLKTFSEKLNLFVESAQTRKIAIRIGISYOGKADNAN 240
      SLLLN+KV+M L N EL +VKGRL KTF+KMLD+++ R IAEI ISY G+A +A
30  Sbjct: 185  SLLLNKIVMALKDELKTLVKGKGNKTFKMLDLYLAKNSHPIAIAISYAGEASLAL 244

Query: 241  NFRKGLAV-LGAPISVLETSIIQTHTGEGAFVMMVRYE 278
      +E++A ISVLETSIIQTHTGE AFVMMVRYE
35  Sbjct: 245  TLKERIAAYNHSISVLETSIIQTHTGEGAFVMMVRYE 283

```

SEQ ID 300 (GBS113) was expressed in *E.coli* as a His-fusion product. Purified protein is shown in Figure 201, lane 8.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 90

A DNA sequence (GBSx0092) was identified in *S.galactiae* <SEQ ID 307> which encodes the amino acid sequence <SEQ ID 308>. Analysis of this protein sequence reveals the following:

```

55  Possible site: 28

    >>> Seems to have a cleavable N-term signal seq.

    ----- Final Results -----
        bacterial outside --- Certainty=0.3000(Affirmative) < succ>
        bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
        bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

55  >GP:CAA72097 GB:Y11213 hypothetical protein [Streptococcus thermophilus]
    Identities = 75/185 (40%), Positives = 116/185 (62%), Gaps = 3/185 (1%)

Query: 13  WKWAFLLALLAINLSPFVAVIASRLIQVREPNTGKISTGVQDKVKVGTFTTNKSQLNKIAL 72

```

-159-

WKW FL LLA+NL+ +V+ R++ E + + G K+G ++ +K +L++++
 Sbjct: 5 WKWFLGLALAINLALISVVTVRIMTPVETSPVSLPEGA---TKIGKYSMSKEELDESIRG 61
 Query: 73 YLKQYQTKRMNYKIYAASSSILFBSGYQLLGYEVPLIYIFRPYRITNGAVQLKVTFSVG 132
 + + Y T KM +K+ +S I+FE SY++LG+ VPLY+YF P +GAV L+ + S G
 Sbjct: 62 FAQDYSTDKMRFKVKVINSKIVPESSTKVLGHAIVPLVYVTFPLVSESGAVVLQESLSAG 121
 Query: 133 TLPLPEKDVLYQLKSSYKLPNFVDIKPKKSVININQLDKNKRGYIKATAIDLNDNFS 192
 TL LP D L IK S KLP+++ I KK + +N+Q +KN +GI +A + DLVND
 Sbjct: 122 TLKLPILDALNMIKRSTKLPDYIVIDSKKGKVIINIQSMKNDKGITARAQSPDLVNDRE 181
 Query: 193 FDIK 197
 FDI+K
 Sbjct: 182 FDIYK 186

A related DNA sequence was identified in *Spyogenes* <SEQ ID 309> which encodes the amino acid sequence <SEQ ID 310>. Analysis of this protein sequence reveals the following:

Possible site: 29

>>> Seems to have a cleavable N-term signal seq.

----- Final Results -----
 bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP:CAW72097 GB:Y11213 hypothetical protein [Streptococcus thermophilus]
 Identities = 73/185 (39%), Positives = 112/185 (60%), Gaps = 3/185 (1%)
 Query: 10 WKWSFLCLLAPNTAFILMVIASRLIQVREPESELIACKPVNIKIGITFVTTRBOLNETVAS 69
 WKW FL LLA N A + V+ R++ E + K K IG + ++E+L+E++
 Sbjct: 5 WKWFLGLLAINLALISVVTVRIMTPVETSPVSLPKGATK---IGKYSMSKEELDESIRG 61
 Query: 70 YLKDYQTEKMSYKFATSSSSILFBSGYQLLGYEVPLIYIFPQHRLENGAVQLQVIFSVSG 129
 + +DY T+KM +K T+S I+FE +Y++LG+ VPLY+YF P E+GAV LQ S G
 Sbjct: 62 FAQDYSTDKMRFKVKVINSKIVPESSTKVLGHAIVPLVYVTFPLVSESGAVVLQESLSAG 121
 Query: 130 TLPLPEKDVLYQLKSSYKLPNFVDIKPKKSVININQLDQNDKAVYLKAKKIDLFNDEIS 189
 TL LP D L +K S KLP + + + +N+Q ++ND + +A+ DL ND
 Sbjct: 122 TLKLPILDALNMIKRSTKLPDYIVIDSKKGKVIINIQSMKNDKGITARAQSPDLVNDRE 181
 Query: 190 FNIYK 194
 F+IYK
 Sbjct: 182 FDIYK 186

An alignment of the GAS and GBS proteins is shown below:

Identities = 129/194 (66%), Positives = 155/194 (79%)
 Query: 5 KIGRNLNFWKWFLLLA LLA LNSFTAVIASRLIQVREPNTGKISGTGVQDKVKGVTFTTNKS 64
 K NLA+WKW+FL LLA N +F VIASRLIQVREP + I+ +K+GTF T +
 Sbjct: 2 KKKSNLNMWWSFLCLLAPNTAFILMVIASRLIQVREPESELIACKPVNIKIGITFVTTRR 61
 Query: 65 QLNKTIALYLKDYQYTKRMNYKIYAASSSILFBSGYQLLGYEVPLIYIFRPYRITNGAVQL 124
 QLN+T+A YLK YQT+KM+YK YA SSSILFBSGYQLLGYEVPLIYIFP+RL NGAVAL
 Sbjct: 62 QLNETVASYLKDYQYTKRMNYKFATSSSSILFBSGYQLLGYEVPLIYIFPQHRLENGAVQL 121
 Query: 125 KVTFSVSGTTLPLPEKDVLYQLKSSYKLPNFVDIKPKKSVININQLDKNKRGYIKATAIDLNDNFS 184
 +V SFSVSGTTLPLPEKDVLYQLKSSYKLP+V + P +S I +NLQD++N +YLGK I
 Sbjct: 122 QVIFSFSVGTTLPLPEKDVLYQLKSSYKLPNFVDIKPKKSVININQLDQNDKAVYLKAKKI 181
 Query: 185 DLVNDNFSFDIFK 198
 DL ND SF+I+KK
 Sbjct: 182 DLFNDEISFNIYK 195

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

A related GBS gene <SEQ ID 8487> and protein <SEQ ID 8488> were also identified. Analysis of this protein sequence reveals the following:

```

Lipop: Possible site: -1  Crend: 7
McG: Discrim Score: 7.47
GVH: Signal Score (-7.5): 2.42
    Possible site: 28
>>> Seems to have a cleavable N-term signal seq.
ALOM program count: 0 value: 5.89 threshold: 0.0
    PERIPHERAL Likelihood = 5.89 120
    modified ALOM score: -1.68

*** Reasoning Step: 3

----- Final Results -----
    bacterial outside --- Certainty=0.3000(Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
    bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

SEQ ID 308 (GBS20) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 4 (lane 5; MW 25kDa) and in Figure 167 (lane 12-14; MW 37kDa – thioredoxin fusion). It was also expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 9 (lane 7; MW 47.6kDa). Purified Thio-GBS20-His is shown in Figure 244, lane 12.

Example 91

A DNA sequence (GBSx0093) was identified in *S.agalactiae* <SEQ ID 311> which encodes the amino acid sequence <SEQ ID 312>. This protein is predicted to be histone-like DNA-binding protein. Analysis of this protein sequence reveals the following:

```

Possible site: 40

>>> Seems to have no N-terminal signal sequence

----- Final Results -----
    bacterial cytoplasm --- Certainty=0.2768(Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
    bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 9313> which encodes amino acid sequence <SEQ ID 9314> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AAD40810 GB:I40355 histone-like DNA-binding protein [Streptococcus mutans]
Identities = 43/47 (91%), Positives = 46/47 (97%)

Query: 1  YANKQQLIAKVARATELTKKDSAAAVDAVFAAVADYLAEGKEKVLIG 47
          YANKQQLIAKVARATELTKKDSAAAVDAVFAAVADYLAEGKEKVLIG
Sbjct: 1  YANKQQLIAKVARATELTKKDSAAAVDAVFAAVADYLAEGKEKVLIG 47

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 313> which encodes the amino acid sequence <SEQ ID 314>. Analysis of this protein sequence reveals the following:

```

Possible site: 25

```

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

5 bacterial cytoplasm --- Certainty=0.2834 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 41/47 (87%), Positives = 44/47 (93%)

10

Query: 1 MANKQDLIAKVAEATELTKKDSAAAVDAVFAAVADYLAEGEKVQLIG 47
 MANKQDLIAKVAEATELTKKDSAAAVDAVF+ + LAEGEKVQLIG
Sbjct: 1 MANKQDLIAKVAEATELTKKDSAAAVDAVFETIRAFLEAGSEKVLIG 47

15

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 92

A DNA sequence (GBSx0094) was identified in *S. agalactiae* <SEQ ID 315> which encodes the amino acid sequence <SEQ ID 316>. Analysis of this protein sequence reveals the following:

20

Possible site: 54

>>> Seems to have no N-terminal signal sequence

25

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2722 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 9293> which encodes amino acid sequence <SEQ ID 9294> was also identified. A further related GBS nucleic acid sequence <SEQ ID 10793> which encodes amino acid sequence <SEQ ID 10794> was also identified.

30

The protein has homology with the following sequences in the GENPEPT database:

>GP:AA17886 GB:AF100456 hyaluronate-associated protein precursor
[Streptococcus equi]

35

Identities = 303/435 (69%), Positives = 360/435 (82%), Gaps = 1/435 (0%)

Query: 1 MATKVDVEKDGLTYTATLRGLKWSGSGLTAKDPVYSWQRLVDPITASQYAYLAVEGHV 60
+A KVDVB+DGLTYTATLR GLKWSGSG LTA+DFVYSWQR+VDPITAS+YAYLA E H+
Sbjct: 87 LAEKVDVEEDGLTYTATLRDGLKWSGSGDLTAEDFVYSWQRMVDPIKASEYAYLATSEHL 146

40

Query: 61 LNADKINGEQKDKLNKLGVKAEGDDKVITLSSPSQFIYLLAFINFMQKQVEVKEYGK 120
NA+ IN G+ DL+ LGVKA+G+ KV+ TL+ P+DQF L+P+NF+PQR+ V+ GK
Sbjct: 147 KNAEDINSGKNFDLDELGVKADGN-KVIFLTETAPQPKSLLSPSNFVQKESFVKDAGK 205

45

Query: 121 DYATTSKIFVYSGPYTVEGWSNGIFTLKKNKNYMDAKIVKTKVRIQTIVKKEPTAVQM 180
DY TTS+ +YSGFY V+ WNG++GTF L KKNKNYMDAKIVK+ V +QTVKKEPTAVQM
Sbjct: 206 DYGTTSEKQIYSGFYIVKDWNGTSGTFKLKKNKNYMDAKIVKTTETVNVQTVKKEPTAVQM 265

50

Query: 181 YKRGELDAANISNTSAIYQANKNNKVDIVLEATTAYMEYNTTSGKGLDNVIRRALNL 240
YK+G+LD ANIS TSAIY ANK +KDV VLEATTAY+ YN TG+++GL+++KIR+ALNL
Sbjct: 266 YKRGELDFANISGTSIAIYNANKKHVDVVFVLEATTAYIVYNGTALIEGLNLSKIRQALNL 325

55

Query: 241 ATNRKGVVQAAVDVTSKPAIAFAPTGLAKTPDGTDLAKYVAPGYENKTEAKLPKEGLA 300
AT+RNG+V AAVDTGSKPA A FTGLAK DGTDL ++VAPGY+Y EAALPKPEGLA
Sbjct: 326 ATDRKGVISAVDVTSKPAIALVFTGLAKLSGDTDLTSHVAPGYKYDKEAKLPKEGLA 385

Query: 301 ESLGTLKLITADADAPAAKNSVDYIKSTWEAALPGLTVEKPVTFKQRLSDSRKQND 360
E G L +TITADADAPAAK++VDYIK TWE ALPGLTVEKPV FKRLED++ QNF+

Sbjct: 386 ELGKDALITITADADAPAAKSAVDYIKETWETALGLIVREKVFVFFKORLETRKQNF 445

Query: 361 IVVSLWGGDYPEGSTFYGLFKSDSQNDGKFKANKDYDAAYNKALISEDAMKPAKSAKYR 420
+ V LMGSDYF+GSTFYGLFKS S N GKF N DYDAAYNKA++ DA+ +A DYK

Sbjct: 446 VAVVWGGDYPEGSTFYGLFKSGSAYNYGKFTNADYDAAYNKALITTDALNTDAADYKA 505

Query: 421 AEKILFEQGYNPLY 435

AEK L++ YNPLY

Sbjct: 506 AEKALYDALNPLY 520

A related GBS gene <SEQ ID 8489> and protein <SEQ ID 8490> were also identified. Analysis of this protein sequence reveals the following:

Lipop: Possible site: 21 Crend: 4

Sequence Pattern: CGSR

SRCLFG: 0

McG: Length of UR: 19

Peak Value of UR: 2.34

Net Charge of CR: 3

McG: Discrim Score: 5.94

GvH: Signal Score (-7.5): 0.6

Possible site: 20

>>> May be a lipoprotein

Amino Acid Composition: calculated from 22

ALOM program count: 0 value: 5.14 threshold: 0.0

PERIPHERAL Likelihood = 5.14 166

modified ALOM score: -1.53

*** Reasoning Step: 3

----- Final Results -----

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GF|4336671|gb|AAD17886.1||AF100456 hyaluronate-associated protein
precursor {Streptococcus equi}

Score = 721 bits (1640), Expect = 0.0

Identities = 354/515 (68%), Positives = 417/515 (80%), Gaps = 2/515 (0%)

Query: 1 KNWRRVGVGLTLASVATLAACGSK-SASQDSNGAINWAIPTETFLDLKVTDTYSNLA 59
K +R+G+ +TLASVA L ACJ+K SAS D INW PTEI TLD+SK TDTYS LA

Sbjct: 7 KACKRLGLAAVTLASVAALMACGNKQASSTDKKSEINWYTPTEITLDTSKNTUTYSALA 66

Query: 60 IGNSSSNFLRLDKDGRPOLATKVDVSKDGLTYTATLRKGLKWSGSKLTAQDFVYSNQ 119

IGNS SN LR D GK +PDLA KVDVS+DGLTYTATLR GLKWSGDS LTA+DFVYSNQ

Sbjct: 67 IGNSGNNLRADAKGLQDLPAEKVDVSDGLTYTATLRGLKWSGSDLTAEQDFVYSNQ 126

Query: 120 RLVDPKTASCYAYLAVRSHVLNADKINEQEKDLNKLGVKABGDDKVVITLSSPSQPIFY 179

R+VDPKTAS+YAYLA E H+ NA+ IN G+ DL+ LGVKA+G+ KV+ TL+ P+POF

Sbjct: 127 RMVDPKTASAYAYLATRSHLNADINSKNPDLDSLGVKADGN-KVIPTLEPAQPFKS 185

Query: 180 YLAFINFMPOKQEVVSKYKDYATTSKNTVYSGPYTVGWSGNGFTFLKNNKNTWDAKN 239

L+P+NF+PK+ V+ GRDY TTS+ +YSGPY V+ WNG++GTF L KNNKNTWDAKN

Sbjct: 186 LLSFSNFMPOKQEVVSKYKDYATTSKNTVYSGPYTVGWSGNGFTFLKNNKNTWDAKN 245

Query: 240 VKTEKVRIGTVKKPDTAVQMYKRGELDANISNTSAIYQANKNNKDVTVLEATTAIYMEY 299

VKT+ V +QTVKKPDTAVQMYK+G+LD ANIS TSAIY ANK +KDV VLEATTAIY+ Y

Sbjct: 246 VKTETVNVQTVKKPDTAVQMYKQGLDLPANISNTSAIYQANKNNKDVTVLEATTAIYV 305

Query: 300 NTTGSVRGLDNVKIRRALNLAIRKGVVQAANDTGSKPALAFAPTGLAKTPDGLAKYV 359

N TG+++GL+++KIR+ALNIAT+RKG+V AANDTGSKEP A PTGLAK DGTDL+ +V+

Sbjct: 306 NQTGAIRGLNLSKIRQANLAIATDRKGVISAAVDTGSKPATALVPTGLAKLSDGTDLTEHV 365

Query: 360 APGYEYNKEAAKLFKQGLASSGLTKLKLTTTADADAPAAKNSVDYIKSTWEAALPGLTV 419
 APGY-Y+ KAALFKQGLAS G L +TTTADADAPAAK++VDYIK TWE ALPGLTV
 Sbjct: 366 APGYKYDDEAAKLFKQGLAELGKDALTTTADADAPAAKNSVDYIKETWETALPGLTV 425

5 Query: 420 EEKFTVIFKQRLIEDSRKONFDIVSLMGSDYPGSGTFYGLFKSDSQNDGKPFANKDYDAAY 479
 EEKFTV FQRLIED++ QNF++ V LMGGDYP-GSTFYGLFKS S N GKF N DYDAAY
 Sbjct: 426 EEKFTVPFKQRLIEDTKNPFVAVLMGGDYPGSGTFYGLFKSGSAYNYGKPFINADYDAAY 485

Query: 480 NKAISEDAMKPAESAKDYKEAKILFSQDAYNPLY 514
 NKA++ DA+ +A DYK ARK L++ YNPLY
 10 Sbjct: 486 NKALTDAINTDAADDYKAARKALYNALYNPLY 520

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 317> which encodes the amino acid sequence <SEQ ID 318>. Analysis of this protein sequence reveals the following:

15 Possible site: 24

>>> May be a lipoprotein

----- Final Results -----
 20 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

25 Identities = 114/428 (26%), Positives = 185/428 (42%), Gaps = 63/428 (14%)

Query: 7 VSKDGLTYTTLIRKGLK--SDGSK---LTAKDFVYSWQRLVDPKTAQYAYLAVEGHVL 61
 VSKDGLTYT TLR G+ W +DG+ +TA+DFV + VD K+ + Y VE +
 30 Sbjct: 92 VSKDGLTYTTLRGVSMYTTADGEGYAPVTAEDFVTLGKHVDDKSDALY---VWEDSIK 148

Query: 62 NADKINEGQEKDLNKLGVKAEGDDKVVITLSSPSQPIYYLAFTNFMQKQVEVKEYGKD 121
 N G E D ++GVKA D V TL+ P + ++ P + ++ GKD
 Sbjct: 149 NLKAYQNG-EVDFKIEVGKALDDTKVQYTLNKPESYWNKSTTYSVLFPVNAKFLKSGKD 207

35 Query: 122 YATTSKNTV-YSGPYTVBWNQSGNSGTFLKKNQFYDAKQNVKTEVRI--QTWKPKDTAV 178
 + TT +++ +G Y + + S + KN+NYWDAQIV + V++ P +
 Sbjct: 208 FOTTDPSSILNGAYFLSAFT-SKSSVEPHKNENFYDAKQNVGIESVLTYSGSDGSPGSPY 266

40 Query: 179 QMYERGELDAANISNTSALYQANKIN--KQVT-DVLEATAYMEYNT----- 223
 + + +GE A + Y++ K N ++T +L ++ N
 Sbjct: 267 KNFKGEPFVARLYPNDFYKSAIGNYADNTITYGMLTGDRIHLTVNLRNTPFNKTKDPA 326

Query: 224 ---GSVKGLDNVKKIRALNLTNRIGVVAQAVDTGSKPA---IAPAPT--GLAKTPDGT 274
 K L+N R++ A +R +K + PT ++ G+
 45 Sbjct: 327 QDAGKKALNNDKFRQAIQAFDRASPAQTAGQDAKTALRNMLVPTTFVTGSDGSGS 386

Query: 275 DLAKYVAP-GYR-----YKTRAAKLF--KEGLASGLT-KLKLTTTADAD 316
 ++ K +A G E YN +A F KB L G+T ++L D
 50 Sbjct: 387 EVEKEAKLGDGWDKVNLDAAQGGFYNPEKAKAEFAKAEALTAGVTFPQGLDVPDQA 446

Query: 317 APAAKNSVDYIKSTWEAALGLTV----EEKFTVIFKQ--LIEDSRKONFDIVSLMG 368
 A K + EA+L V E + T + + E +Q++DI+ S WG
 Sbjct: 447 NAATVQEAQSFQSGVEASLGKENVIVNVLKTRISTHEAQGFYAEFTPQDQYDISSWVG 506

55 Query: 369 DYPEGSGTF 376
 DY + T+
 Sbjct: 507 DYQDPRTY 514

SEQ ID 9294 (GBS663) was expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 137 (lane 3; MW 89.5kDa). It was also expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 137 (lane 5-7; MW 64.5kDa), in Figure

179 (lane 11; MW 65kDa) and in Figure 65 (lane 2; MW 61kDa). Purified GBS663-His is shown in Figure 231, lane 3-4. Purified GBS324-His is shown in lane 6 of Figure 210.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

5 Example 93

A DNA sequence (GBSx0095) was identified in *S.agalactiae* <SEQ ID 319> which encodes the amino acid sequence <SEQ ID 320>. This protein is predicted to be transmembrane protein OppB (oppB). Analysis of this protein sequence reveals the following:

```

Possible site: 37
10  >>> Seems to have no N-terminal signal sequence
    INTEGRAL    Likelihood = -10.77    Transmembrane    293 - 309 ( 281 - 313)
    INTEGRAL    Likelihood = -9.77    Transmembrane    21 - 37 ( 14 - 46)
    INTEGRAL    Likelihood = -6.32    Transmembrane    115 - 131 ( 105 - 132)
15  INTEGRAL    Likelihood = -4.88    Transmembrane    144 - 160 ( 140 - 166)
    INTEGRAL    Likelihood = -3.03    Transmembrane    238 - 254 ( 237 - 255)

---- Final Results ----
20  bacterial membrane --- Certainty=0.5310 (Affirmative) < succ>
    bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
    bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 8491> which encodes amino acid sequence <SEQ ID 8492> was also identified.

25 The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AAF73091 GB:AF103793 transmembrane protein OppB [Listeria monocytogenes]
Identities = 147/304 (48%), Positives = 221/304 (72%), Gaps = 1/304 (0%)

Query: 13 MIKYILKRVAILLVTLWVVITLSFFLMQILPGTTPYNNIP-KLTREMIALLNKQYGLDKPWN 71
30  M+KY LKRV +L+TL+++ ++F LM+ LPGTPY N KL+E I + N++YGL+ +
Sbjct: 1 MVKYTLKRVLNMLITLFIILASVTFFVMKPLPGTPYRNQKLSDEQIMHMKYGLNDISIP 60

Query: 72 QQVLYTLNVLHGDFTSYQSVNPQVSRMISLRIGSVSLGVQALVFGVLGGILVGAISA 131
35  QY Y+ ++ GD G S+Q N+PVS ++S +G SV L ++A+ FG+ GIL+G I+A
Sbjct: 61 VQYFNMYMTGLVKGDIAGVSQOLDNRPVSEILSALIGPSVQLALEAMAFGVIFGILLGVIA 120

Query: 132 RHKNDKVDGILSVIATLGLISMPSPFIIGILLLLDYFGFKNLLPLSGWGTFSQTILPSLAL 191
40  ++N D + IA LG S+PSF+ +L + G K + P++GWTG+ TILP+ AL
Sbjct: 121 MYQNRWPDTSTFIILGKSVSPFVFATVLYQLGAKLQIFPVAGWGTGADTILPAFALA 180

Query: 192 LPTLASVSRRFSRSHMIETLNSDYQVARSKGMITIRQVTRKHAYRNSMIPILTLIQLAAG 251
45  + LA+ +RF R+E+I+ SDTV LA++KG + +V KHA RN++IP++T++GPL+
Sbjct: 181 MFPLATAARFMTETLIDVFPASDYLILAKAGNSRTIEVAVKHAIENALPLITVLGSLVA 240

Query: 252 LITGSALIEQIFSIPIGQQQFVTSIPTKDYPIVMGTTITVAVMLVAAILLDDVVISIVDP 311
50  L+TGS +IE I+SPIGIG QFV+SI T DYPVIMGTTI++AVML+ IL+ D++ ++DP
Sbjct: 241 LMTGSVLENIYSIPIGSGFVSSIQTNDYPIVMGTTITLFAVMLVFVLWVDVLYGLIDP 300

Query: 312 RVRL 315
50  R+R+
Sbjct: 301 RIRV 304

```

There is also homology to SEQ ID 64.

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 9069> which encodes amino acid sequence <SEQ ID 9070>. Analysis of this protein sequence reveals the following:

-165-

Possible site: 25

>>> Seems to have an uncleavable N-term signal seq

5	INTEGRAL	Likelihood = -8.81	Transmembrane	466 - 482 (463 - 493)
	INTEGRAL	Likelihood = -5.10	Transmembrane	419 - 435 (418 - 440)
	INTEGRAL	Likelihood = -4.78	Transmembrane	328 - 344 (322 - 348)
	INTEGRAL	Likelihood = -4.41	Transmembrane	366 - 382 (365 - 384)
	INTEGRAL	Likelihood = -4.09	Transmembrane	290 - 306 (287 - 311)
	INTEGRAL	Likelihood = -2.97	Transmembrane	17 - 33 (13 - 36)

10 ----- Final Results -----

bacterial membrane --- Certainty=0.4524 (Affirmative) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

15 An alignment of the GAS and GBS sequences follows:

Score = 117 bits (291), Expect = 3e-28

Identities = 61/208 (29%), Positives = 121/208 (57%), Gaps = 4/208 (1%)

20 Query: 291 IGPFVGFVMSIVGLPLGLFMAFKNYTFDSFSTATIMFMLALPSIAV-IYVVRFLGGMWG 349

+G ++P +G +G AR KN D + T +++S + I ++ + G

Subject: 99 LGVQALVFGVLGGILVGAISARKKNDKVDGILSVIATLGISWPSFIIGILLDYFGPKKN 158

Query: 350 LPDSFPMGLGRSDPKSYILPALILGILNIPTTVIMFRKYAVDLQASDMVRFARSKGLSESE 409

L P+G ILP+L LG+ + + FR ++ SD+V+ ARSKG+ +

25 Subject: 159 L---LPLSGWGTFSQVITLPSIALGLPTLASVSRFRSEMITLNSDYQLARSKMTIRQ 215

Query: 410 IYRHLFKNAMVPTVSGVPASIIAIGGATLTETVFAPFGMGKMLDSIKSANNMIVGL 469

+ R H ++N+M+PI++ + +G+ L E +F+ PG+G+ + SI + + +I+G

30 Subject: 216 VTRKHAYRNSMIPILTLIGPLAAGLLTGSALIEQIFSIPIGIGQFVTSIPTKDYFVIMST 275

Query: 470 TFIPTVLSIVSLILGDIWNTLVDPRIKL 497

T ++ V+ +V+L+ D+V++VDPRL+L

Subject: 276 TIVAVNMLVAILITDVVISIVDPVRL 303

35 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 94

A DNA sequence (GBSx0096) was identified in *S. agalactiae* <SEQ ID 321> which encodes the amino acid sequence <SEQ ID 322>. This protein is predicted to be transmembrane protein OppC (oppC). Analysis of

40 this protein sequence reveals the following:

Possible site: 59

>>> Seems to have no N-terminal signal sequence

45	INTEGRAL	Likelihood = -11.52	Transmembrane	311 - 327 (307 - 333)
	INTEGRAL	Likelihood = -7.80	Transmembrane	42 - 58 (40 - 65)
	INTEGRAL	Likelihood = -7.43	Transmembrane	142 - 158 (131 - 165)
	INTEGRAL	Likelihood = -4.73	Transmembrane	182 - 198 (179 - 214)
	INTEGRAL	Likelihood = -3.50	Transmembrane	257 - 273 (257 - 276)

50 ----- Final Results -----

bacterial membrane --- Certainty=0.5607 (Affirmative) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

55 The protein has homology with the following sequences in the GENPEPT database:

>GP:AAF73092 GB:AF103793 transmembrane protein OppC [Listeria monocytogenes]

Identities = 157/325 (48%), Positives = 219/325 (67%), Gaps = 4/325 (1%)

60 Query: 20 EKIEKPALSFMDNRRLKKNKLAVVSLYLALLLTFSASNLFPVTRKANGFDSKKVIT 79

-166-

```

      EKI +P+L+P+QD+W R++KNK A+VSL +IAL++ ++          ++++T
Sbjct: 22 EKINRPSLTPLQDSWLIRKKNAAVSLVLVAIVTMAIVGVPLSCLNGPENNINRQITE 81

5  Query: 80 YRNLPPKLS--NLFFWNGSIKYAGNTESTDAYKSONVPEKVKALGTDLSGRSVAKRI 137
      +LPPK+  N+PFWNG  G R D YK N+ R Y LG+D+LGR RI
Sbjct: 82 NASLPEKVGQGFENPFWNGHSIQG--RDVDIYKQNNIKEGTYIWLQSDTIGRDQFARIW 139

      QVIRISLVAIAATFIDLIIGVTYGLVSGFAGGRLDTIMQRIVEVSIIPNLVIVTMLGL 197
      G R+SL++A+ A DL+IGV YGL+SG+ GGR+D MQR++EVI +IPNLV++ V + L
10 Sbjct: 140 AGTRVSLIIAVVAALCDLVIGVAYGLISGVVGRVDNFMQRVLEVIQATPNLNVIVTML 199

      VLNGITAIISIAFTGWTSMSRQVNMILSYREREFVLAARSLGESPIKIAFKHILPNI 257
      +L GI +III+IA T W +M+R VR L + +RFV+A+ +LGES KI KH+PNI
15 Sbjct: 200 ILEPGIVSIITATMTSWITMARVVROQLKRNQEFVMASTLIGBSTPKILIKHILPNI 259

      SGIIIVQIMTIPSAIMYEAVLSAINLGVPKPTASLGSLSIDQENLQYYPQVILPALA 317
      SGIII+ IM +IPSAI +EA LS I LG+ P ASLG L++D + LQ PY ++ P +
Sbjct: 260 SGIIINIMFSPSAIFPEAFLSFIGLGLPAPASLGLVNDGYKTLQVLPMYILPCIV 319

20 Query: 318 LVMSISLAFILLGDLRDAFDPKSS 342
      L +I +AF L+ DGLRDAFDPK D
Sbjct: 320 LCIIMIAFNLIADGLRDAFDPKMRD 344

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 323> which encodes the amino acid sequence <SEQ ID 324>. Analysis of this protein sequence reveals the following:

Possible site: 59

```

>>> Seems to have no N-terminal signal sequence
30 INTEGRAL Likelihood = -10.30 Transmembrane 43 - 59 ( 37 - 65)
INTEGRAL Likelihood = -8.49 Transmembrane 111 - 127 ( 109 - 135)
INTEGRAL Likelihood = -6.26 Transmembrane 279 - 295 ( 270 - 298)
INTEGRAL Likelihood = -3.88 Transmembrane 172 - 188 ( 172 - 188)
INTEGRAL Likelihood = -3.61 Transmembrane 145 - 161 ( 145 - 165)
35 INTEGRAL Likelihood = -1.49 Transmembrane 223 - 239 ( 223 - 239)

----- Final Results -----
      bacterial membrane --- Certainty=0.5118 (Affirmative) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
40      bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

```

Identities = 91/325 (28%), Positives = 156/325 (48%), Gaps = 34/325 (10%)

45 Query: 16' SSTQEKIEKPAISFMQDAWRRLKKNKLAVVSLYLLALLTFSLASNLFTVQDKANGFDSK 75
      S E I+ PA S+ + +R+ K V L +L +L S +F +D
Sbjct: 16 SEASEVIDTPAYSYSKSVPROFFSKKSTVFMVLVILVTVMMSFIYMFAN-----YDFN 69

      KVITYRNLPPKLSNLPFWNGSIKYAGNTESTDAYKSONVPEKVKALGTDLSGRSVAKR 135
      V+ + + + + +Y GTD G+S+
50 Sbjct: 70 DVSNIND-----PSKRYINENAEYFGTDKNQGSFLDG 102

      IIVGIRISLVAIAATFIDLIIGVTYGLVSGFAGGRLDTIMQRIVEVSIIPNLVIVTML 195
      + G R S+L++A+ AT I++ IGV G + G + D +M I +IS+IP+++I+ L
55 Sbjct: 103 VYTGARNISILISVIATLINITIGVVLGAIMGVSKA-FDKVMIEIYNIISNIPMLIIIVL 161

      GLVLNGITAIISIAFTGWTSMSRQVNMILSYREREFVLAARSLGESPIKIAFKHILP 255
      LG G +I++ TGW ++ +R L YR+ E+ LA++LG KIA K++LP
Sbjct: 162 TYSLGAGFNWLIACFITGWIGVAYSIRVOILLRYRDLSEYLNASCTLGTMPYKIAVKNILP 221

60 Query: 256 NISGIIIVQIMTIPSAIMYEAVLSAINLGVPKPTASLGSLSIDQENLQYYPQVILPALA 315
      + +I+ + +P + EA LS +G+ T SLG I++ NL Y +P
Sbjct: 222 QLVSVIMTMLSQMLQSVVYSSEAFLSFGFGLPTTTPSLGRFTANYSSNLITNAYLFWIPL 281

      IALVMSISLAFILLGDLRDAFDPKS 340
65 + L+++SL ++G L DA DP+S

```

Sbjct: 282 VTILVLSPLLYVGNLADSDPRS 306

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

5 Example 95

A DNA sequence (GBSx0097) was identified in *S. agalactiae* <SEQ ID 325> which encodes the amino acid sequence <SEQ ID 326>. This protein is predicted to be ATPase OppD (oppD). Analysis of this protein sequence reveals the following:

Possible site: 20

>>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.85 Transmembrane 164 - 180 (163 - 180)

----- Final Results -----

bacterial membrane --- Certainty=0.1341(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAF73093 GB:AF103793 ATPase OppD [*Listeria monocytogenes*]

Identities = 230/342 (67%), Positives = 283/342 (82%), Gaps = 2/342 (0%)

Query: 4 ETILSVNHLHVDPHFYTAGEVKAIKRVNFKKGETLAIVGESGSGKSVTTTTLGLNAK- 62
 E +L V +L++ PHYTAGEVKAIKRVNFKKGETLAIVGESGSGKSVTT++L +
 Sbjct: 2 EKLEVKLNLNISFHTYTAGEVKAIKRVNFKKGETLAIVGESGSGKSVTTKIMLLPEG 61

Query: 63 NSEI-SGNVQFQGRNLVELSEBWTKVRNGEISMIPODPMTSLDPTMKIGMOIAEPMH 121
 NSEI SG + F G ++ + E++ K+RG +I+MIFQDPMTSL+PTM IG QI+EP++ H
 Sbjct: 62 NSEIKSGQLLPHGMDIAKHEKCMQKIRGKDIAIMIPODPMTSLMPTMTIGKQISEPLIKH 121

Query: 122 QKISKDKALKLALMLKDVGINASEHINDYPHWSGGMRQRAVIAIALAADPEILLIADE 181
 QKISK +A K AL L++ VGI NAE I YPHQ+SGGMRQRAVIA+LA +P+ILLIADE
 Sbjct: 122 QKISKHEAHKIALRLQLVGIANAERI KYPHQSFGGMRQRAVIAISLACNPOLLIADE 181

Query: 182 PTTALDVTIQAILNLKCKIAQRDSSIVFTIHDGLGVVAGMADRAVVMYAGKIVEFGTVD 241
 PTTALDVTIQAIL+LKK +Q + D+SI+FTIHDGLGVV +ADRAVVMY GIVE GTVD
 Sbjct: 182 PTTALDVTIQAILDLKDLQKIDTSIIFTHDGLGVVADRAVVMYGGKIVEFGTVD 241

Query: 242 EVFYNQPHFYTWGLNSMPTTDTESGSLBSIPGTPDLLNPPKGDAPARNFALDIDHE 301
 E+FYNQPHFYTWGL+SMPT DT+ L IGPDPDLL+PPKGDAPARN++A+ ID E
 Sbjct: 242 EBFYNQPHFYTWGLISSMPTLTDDEELVPIGTPDLLHPPKGDAPARNKYMQIDLE 301

Query: 302 BEPPYFKVSETHFAATWLLDERSPKVLPPLPIQKRMERWNEI 343
 BEPP FKVS+TH+ATWLL +P+V PP +R E++ E+
 Sbjct: 302 BEPPLFKVSITHYAATWLLHDAPEVTPDAVLRROEQEAEI 343

There is also homology to SEQ ID 72.

SEQ ID 326 (GBS375) was expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 64 (lane 9; MW 42kDa). It was also expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 71 (lane 3; MW 67kDa).

GBS375-GST was purified as shown in Figure 215, lane 10.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 96

A DNA sequence (GBSx0098) was identified in *S.galactiae* <SEQ ID 327> which encodes the amino acid sequence <SEQ ID 328>. Analysis of this protein sequence reveals the following:

```

Possible site: 28
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3060(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AAA62692 GB:M57689 sporulation protein [Bacillus subtilis]
Identities = 195/308 (63%), Positives = 245/308 (79%), Gaps = 4/308 (1%)

Query: 1 MTENRKLLVEVKNVSLTFNKGKANEVRAIDNVSFDIYBGEVFLGVGESGSGKTTVGRSIL 60
      M E + KL+E+K++ F + V+A+D++SFDIY+GE GLVGRSG GK+T GRSI+
Sbjct: 1 MNELTEKLLLEIKHLKHQHFVTPROT-VCAVDDLSDFIYKGTLLGVGESGCGKSTTGRSI 59

Query: 61 KLYDISDGEITFNGEVISHLKG-KALHSFRKDAQMFQDPQASLNGRMKIRDIVABGLDI 119
      +LY+ +DGE+ FNGE + K K L F + QMIFQDP ASLN RM + DI+ABGLDI
Sbjct: 60 RLYEATDGEVLFNGENVHGRKSRKLLLEPNRKQMIFQDPYASLNPRMTVADIABGLDI 119

Query: 120 HKLAKSKSDRDEKVQALLDLVGLNKDHLTRYPHFSGGQQRIGIARALAVEPKFIAD 179
      HKLAK+K +R +V LL+ VGLNK+H RYPHFSGGQQRIGIARALAV+P+FIAD
Sbjct: 120 HKLAKTKGERMQRVHLELTVGLNKEHANRYPHFSGGQQRIGIARALAVDPFIAD 179

Query: 180 PISALDVSIQAQVNVNMQKLREQGLTYLFAIHLMSVKYISDRIGVMHWKGLLEVGTSD 239
      PISALDVSIQAQVNVNMQKLREQGLTYLFAIHLMSVKYISDRIGVMH+GKL+E+ +D
Sbjct: 180 PISALDVSIQAQVNVNMQKLREQGLTYLFAIHLMSVKYISDRIGVMYFGKLVBLAPAD 239

Query: 240 DVYNNPIHPYTKSLLSAIPEDPESERQVRHQPYNPAIDQ--DQGERQMEHITPGHPVL 297
      ++Y NP+HPYTKSLLSAIP FDP+ ER RV Q Y+P++ Q DG+ + E+ PGHPV+
Sbjct: 240 ELYENPLHPYTKSLLSAIPDPDYENKRVQKYDPSVHQLDGETMEFRVKGHPFVNC 299

Query: 298 TPQEAEEY 305
      T E + +
Sbjct: 300 TEAEFKAF 307

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 329> which encodes the amino acid sequence <SEQ ID 330>. Analysis of this protein sequence reveals the following:

```

Possible site: 47
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3900(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

```

Identities = 164/306 (53%), Positives = 228/306 (73%), Gaps = 3/306 (0%)

Query: 6 KKLVEVKNVSLTFNKGKANEVRAIDNVSFDIYBGEVFLGVGESGSGKTTVGRSILKLYDI 65
      +KLVEVK++ ++F+GK V A+ N +F I +GE F LVGESGSGKTT+GR+I+L D
Sbjct: 3 EKLVEVKLELISFGSGKKFV-AVNANFNFIKGETFSLVGESGSGKTTI+GRTIGLNDT 61

Query: 66 SDGEITFNGEVISHLKGKA-LHSFRKDAQMFQDPQASLNGRMKIRDIVABGLDIHLK 124
      S G+I ++G+VI+ K K+ + + QMIFQDP ASLN R + I++BGL L K
Sbjct: 62 SSGQILYDGKVNKRKSKSEANELIRKIQMFQDPASLNERATVDYIISGLYFNFLPK 121

```

Query: 125 SKSDRDSKVQALLDLVGLNKDELTRYPHFSGGQRQRIQIARALAVEPKFTIADPISAL 184
 ++ +R K++ ++ VGL +HILTRYPHFSGGQRQRIQIARAL + P+F+IADEPISAL
 5 Sbjct: 122 TEERKEKIKNNMAVGLLSEHILTRYPHFSGGQRQRIQIARALVNNPFIADPISAL 181

Query: 185 DVSIAQVAVNLMQKIQREQGLTYLFTAHDLMSVKYISDRIGVMHMGKLEVGSTDDVYNN 244
 DVS++AQV+NL++++Q E+GLTYLFTAHDL+V++ISDRI V+H G ++EV ++++NN
 10 Sbjct: 182 DVSVAQVNLNLMKQMAEKGLTYLFTAHDLGVVRFISDRIAVTHKGVIVEVASTEELNN 241

Query: 245 PIHPYTKSLLSAIPDPESERQRVHQVYNPAIEQDQGER-QMREITTCGHFVLSTPQEA 303
 PIHPYT+SLLSA+P PDP ERQ+ Y+P ++ M RI P HFV + E E
 15 Sbjct: 242 PIHPYTQSLLSAVPIPDILRQKELVVYHPDQHDYTLDKPMSVVEIKPNHFVWNAQAEI 301

Query: 304 EYKQI 309
 +Y+K++
 15 Sbjct: 302 KYQEL 307

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

20 Example 97

A repeated DNA sequence (GBSx0099) was identified in *S.galactiae* <SEQ ID 331> which encodes the amino acid sequence <SEQ ID 332>. Analysis of this protein sequence reveals the following:

Possible site: 28

25 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm	---	Certainty=0.3021(Affirmative)	< succ>
bacterial membrane	---	Certainty=0.0000(Not Clear)	< succ>
bacterial outside	---	Certainty=0.0000(Not Clear)	< succ>

30

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

35 Example 98

A repeated DNA sequence (GBSx0100) was identified in *S.galactiae* <SEQ ID 333> which encodes the amino acid sequence <SEQ ID 334>. Analysis of this protein sequence reveals the following:

Possible site: 24

40 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm	---	Certainty=0.0352(Affirmative)	< succ>
bacterial membrane	---	Certainty=0.0000(Not Clear)	< succ>
bacterial outside	---	Certainty=0.0000(Not Clear)	< succ>

45

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

50 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 99

A repeated DNA sequence (GBSx0101) was identified in *S.agalactiae* <SEQ ID 335> which encodes the amino acid sequence <SEQ ID 336>. Analysis of this protein sequence reveals the following:

```

5      Possible site: 23
      >>> Seems to have no N-terminal signal sequence

      ----- Final Results -----
10      bacterial cytoplasm --- Certainty=0.5857(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

15 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 100

A repeated DNA sequence (GBSx0103) was identified in *S.agalactiae* <SEQ ID 337> which encodes the amino acid sequence <SEQ ID 338>. Analysis of this protein sequence reveals the following:

```

20      Possible site: 14
      >>> Seems to have no N-terminal signal sequence

      ----- Final Results -----
25      bacterial cytoplasm --- Certainty=0.1472(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

30 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 101

35 A repeated DNA sequence (GBSx0104) was identified in *S.agalactiae* <SEQ ID 339> which encodes the amino acid sequence <SEQ ID 340>. Analysis of this protein sequence reveals the following:

```

      Possible site: 13
      >>> Seems to have no N-terminal signal sequence

40      ----- Final Results -----
      bacterial cytoplasm --- Certainty=0.0111(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

45 The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 102

A repeated DNA sequence (GBSx0105) was identified in *S.agalactiae* <SEQ ID 341> which encodes the amino acid sequence <SEQ ID 342>. Analysis of this protein sequence reveals the following:

Possible site: 20

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

```
bacterial cytoplasm --- Certainty=0.5628(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 103

A repeated DNA sequence (GBSx0106) was identified in *S.agalactiae* <SEQ ID 343> which encodes the amino acid sequence <SEQ ID 344>. Analysis of this protein sequence reveals the following:

Possible site: 39

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

```
bacterial cytoplasm --- Certainty=0.2059(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 104

A repeated DNA sequence (GBSx0107) was identified in *S.agalactiae* <SEQ ID 345> which encodes the amino acid sequence <SEQ ID 346>. Analysis of this protein sequence reveals the following:

Possible site: 21

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

```
bacterial cytoplasm --- Certainty=0.2045(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 105

- 5 A DNA sequence (GBSx0108) was identified in *S.galactiae* <SEQ ID 347> which encodes the amino acid sequence <SEQ ID 348>. Analysis of this protein sequence reveals the following:

Possible site: 36

>>> Seems to have no N-terminal signal sequence

10

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3031(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

15

The protein has homology with the following sequences in the GENPEPT database:

>GP: CAB11822 GB: Z99104 similar to hypothetical proteins [Bacillus subtilis]
Identities = 125/282 (44%), Positives = 184/282 (64%)

20

Query: 1 MKIFEKAPAKLNLGLDIEGRCDGYSGLAMIMVSDIMNDYVITISELKEDCIVIDSQSKM 60
M+I EIKAPAK+NL LD+ + DGYHE+ MIM +IDL D + ++EL ED + + S + +
Sbjct: 1 MRILEKAPAKINLSLDVIRKPDGYHVEIMITITDLADRIEL/TELADEVYSSNRFV 60

25

Query: 61 PLNNNDNVFKAADITIKQYQINGKGVHIREKSIIPVCGGLGGSTDAATIRAINRNWNLQ 120
P + N ++AA +IK++Y + KGV I + K IPV AGL GGS+DAAT+R LNRILWNL
Sbjct: 61 PDCQWNLAYQAQKLKDRYVFKGVSIIMTKVLPVAAGLAGSSDAATLRGLNRILWNL 120

30

Query: 121 NDYDEMVAIGFKIGSDVPYCLGGGCSLVLGKGEIVKPLPTLRPCWIVLVKPDFGISTKSI 180
+ + + +G +IGSDV +C+ GG +L G+GE +K + T CW++L KP G+ST +
Sbjct: 121 LSABT+LALGAEIGSDVFCVYGGTALATGRGEKIKHISTPPHCWVILAKPTIGVSTARV 180

35

Query: 181 FRDIDCKSISRVDIDLKSAILLSQVQLMVKMGNSLEDITITKNFVISTIKERMINSQA 240
+R + I D+ + AI +Q M +GN LE+T+ +P ++IK +M GA
Sbjct: 181 YRALKIDGIEHPDVQGMIEAIEBKSFQCMCSRLGNVLSVTLMDHFEVAMIKIQMKRFGA 240

Query: 241 DVALMTGSGPTVFSMCSIEKKADRVFNSMGKFCKEVIKVELL 282
D IM+GSGPTVF + E K R++N ++GFC +VY VR++
Sbjct: 241 DAVIMSGSGPTVFLGVQYESKVRQYIKGLRGFCQVYAVEMT 282

40

- A related DNA sequence was identified in *S.pyogenes* <SEQ ID 349> which encodes the amino acid sequence <SEQ ID 350>. Analysis of this protein sequence reveals the following:

Possible site: 44

>>> Seems to have no N-terminal signal sequence

45

INTEGRAL Likelihood = -2.87 Transmembrane 28 - 44 (27 - 45)

----- Final Results -----

bacterial membrane --- Certainty=0.2147(Affirmative) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

50

An alignment of the GAS and GBS proteins is shown below:

Identities = 33/52 (63%), Positives = 38/52 (72%)

55

Query: 126 NVAIGFKIGSDVPYCLGGGCSLVLGKGEIVKPLPTLRPCWIVLVKPDFGIST 177
M+ IG IGSVDVYCL GC+ V GKGE+V + L W+VLVKPDFGIST
Sbjct: 1 MMDIGIPIGSDVPYCLGGCAQVTKGKGVCRILGLLSSWVVLKPDFGIST 52

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 106

A DNA sequence (GBSx0109) was identified in *S.galactiae* <SEQ ID 351> which encodes the amino acid sequence <SEQ ID 352>. This protein is predicted to be AdcR protein. Analysis of this protein sequence reveals the following:

Possible site: 19

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1264 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAA96184 GB:Z71552 AdcR protein [Streptococcus pneumoniae]
Identities = 77/146 (52%), Positives = 117/146 (79%)

Query: 1 MTVLEQKLDHLVSQLLKARNQHLLFGTCQSDVKLTNTQSHILMLLSQQLNSDLAKK 60
M L + ++ +++++L+RKNQHE+L G C S+V LINTQSHILMLLS+R LENS+LA++
Sbjct: 1 MRQLANDINAFINEVILQARNQHLLIGHCTSEVALNTQSHILMLLSSESLNSLARR 60

Query: 61 LNIQAQAVTKAVKSLISQMLKANKDSKDARITYFELSLAKPIADENTHHNTLVGVG 120
LN+SQRAVTKA+KSL+ + ML+ +RDSKDAR+ +++L+LA+PIA+KH HHH++TL Y
Sbjct: 61 LNVQAQAVTKAIKSLVKEGMLETSDSKDARVIFYQLTLARPIASEHHHHHHHTLLTYE 120

Query: 121 RLNVHFSKDEKVLERFLDLPSRELE 146
++ F+ +R+ V++RFL E++
Sbjct: 121 QVATQPTFENDQKVIQRFLTALVGEEK 146

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 353> which encodes the amino acid sequence <SEQ ID 354>. Analysis of this protein sequence reveals the following:

Possible site: 28

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1536 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 106/147 (72%), Positives = 126/147 (85%)

Query: 1 MTVLEQKLDHLVSQLLKARNQHLLFGTCQSDVKLTNTQSHILMLLSQQLNSDLAKK 60
M +L+KLD+LV+ ILLKARNQHLLFG CQSDVKLTNTQSHILMLLSQ++LTI+DLAK
Sbjct: 1 MGILERKLDREAVNTILLKARNQHLLFGACQSDVKLTNTQSHILMLLSQQLNTVTLAKA 60

Query: 61 LNIQAQAVTKAVKSLISQMLKANKDSKDARITYFELSLAKPIADENTHHNTLVGVG 120
LNIQAQAVTKA+KSL+ QML KD+ DAR+TYFEL+ELAKPIA EHHHH TL VY
Sbjct: 61 LNIQAQAVTKAIKSLVQMLAGTKDTVDARITYFELSLAKPIASEHHHHHHHTLLTYN 120

Query: 121 RLNVHFSKDEKVLERFLDLPSRELE 147
RL+ FS E +++++F+ +F+ ELGG
Sbjct: 121 RLQKFSAKREAVDKFVTVFARLEGG 147

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 107

A DNA sequence (GBSx0110) was identified in *S.galactiae* <SEQ ID 355> which encodes the amino acid sequence <SEQ ID 356>. This protein is predicted to be AdcC protein. Analysis of this protein sequence reveals the following:

Possible site: 43

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1089(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAA96186 GB:Z71552 AdcC protein [Streptococcus pneumoniae]
Identities = 182/231 (78%), Positives = 206/231 (88%)

Query: 1 MRYITVSGLTFCYDSDFVLEGVNHYLDSGEFVLTGNGAAKSTLIKATLGILTPKGV 60
MRYITV L+F YD +FVLE +NY +DSGEFVLTGNGAAK+TLIKA+LGIL P+G V
Sbjct: 1 MRYITVEDLSFYFDKSFVLEHINYCVDSGEFVLTGNGAAKSTLIKASLGILQPRIGKV 60

Query: 61 NISKENKEGKKLRIAYLPQQIASFNAGFPSSVYEFVKSGRYPNGWFRRLYKHDESHIRV 120
ISK N +GKKLRIAYLPQQIASFNAGFPSS+VYEFVKSGRYPNGWFRRL HDESHI+
Sbjct: 61 AISKNTYQKKLRIAYLPQQIASFNAGFPSTVYEFVKSGRYPNGWFRRLAHDESHIKA 120

Query: 121 SLEAVGMWNRHKKIGSLGGQKQRAVIARMFASDDPFI+LDEPTTGMAGTTEKFEYELM 180
SL++VGMW+R K+GSLGGQKQRAVIARMFASDDP+F+LDEPTTGMAG+ +FYELM
Sbjct: 121 SLDVGMWNRHDKRLGSLGGQKQRAVIARMFASDDPFI+LDEPTTGMAGTTEKFEYELM 180

Query: 181 HHSNATHGKSVLMITHDPFEVKGADYADRNHILVRNQLPWRCFNVHTNMEV 231
HH+AH HGK+VLMITHDP+EVK YADRNHILVRNQL PWRCFNVH N EV
Sbjct: 181 HHSNATHGKAVLMITHDPFEVKGADYADRNHILVRNQLPWRCFNVHTNMEV 231

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 357> which encodes the amino acid sequence <SEQ ID 358>. Analysis of this protein sequence reveals the following:

Possible site: 43

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2722(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 190/232 (81%), Positives = 214/232 (91%)

Query: 1 MRYITVSGLTFCYDSDFVLEGVNHYLDSGEFVLTGNGAAKSTLIKATLGILTPKGV 60
MRYI+V L+FCY+S+FVLEG+ YHLDSEFVT+TGNGAAKSTLIKATLGIL PK G V
Sbjct: 1 MRYISVRLSFPYHSEFVLEGIYTHLDSGEFVTMTGNGAAKSTLIKATLGILQPKAGRV 60

Query: 61 NISKENKEGKKLRIAYLPQQIASFNAGFPSSVYEFVKSGRYPNGWFRRLYKHDESHIRV 120
I+K+NR+GK+LRIAYLPQQ+ASFNAGFPSS+VYEFVKSGRYPNGWFRRLKHDESHIRV
Sbjct: 61 TIAKKNKDGKQLRIAYLPQQIASFNAGFPSTVYEFVKSGRYPNGWFRRLKHDESHIRV 120

Query: 121 SLEAVGMWNRHKKIGSLGGQKQRAVIARMFASDDPFI+LDEPTTGMAGTTEKFEYELM 180

```

SLEAVGMW+NRHK+IGLSGGQKQR VIARMFASDFDIFVLDEPTIGMD+GTT+ FYELM
Sbjct: 121 SLEAVGMWENRHKRIGLSGGQKQRVVIARMFASDFDIFVLDEPTIGMDGTTDIFYELM 180

Query: 181 HHNAHKGKSVLMTIDHPDEVKQYADRNHLVRNQSLFWRCFN+H E + E
1 HH+AH+HGKSVLMTIDHP+EVK YADRNHLVRNQ LFWRCFN+H E + E
Sbjct: 181 HHSAHQHGKSVLMTIDHPREVKAYADRNHLVRNQSLFWRCFN+HEATDEE 232

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

10 Example 108

A DNA sequence (GBSx0111) was identified in *S.agalactiae* <SEQ ID 359> which encodes the amino acid sequence <SEQ ID 360>. Analysis of this protein sequence reveals the following:

```

Possible site: 36

>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.2299(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
20 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 109

A DNA sequence (GBSx0112) was identified in *S.agalactiae* <SEQ ID 361> which encodes the amino acid sequence <SEQ ID 362>. This protein is predicted to be AdcB protein (znuB). Analysis of this protein sequence reveals the following:

```

30 Possible site: 36

>>> Seems to have no N-terminal signal sequence

INTEGRAL Likelihood =-14.33 Transmembrane 145 - 161 ( 136 - 172)
INTEGRAL Likelihood =-11.57 Transmembrane 29 - 45 ( 20 - 47)
35 INTEGRAL Likelihood =-10.56 Transmembrane 261 - 277 ( 255 - 280)
INTEGRAL Likelihood =-8.70 Transmembrane 231 - 247 ( 227 - 253)
INTEGRAL Likelihood =-5.63 Transmembrane 101 - 117 ( 99 - 121)
INTEGRAL Likelihood =-4.94 Transmembrane 186 - 202 ( 183 - 225)
INTEGRAL Likelihood =-3.82 Transmembrane 55 - 71 ( 54 - 74)
40 INTEGRAL Likelihood =-3.61 Transmembrane 206 - 222 ( 203 - 225)
INTEGRAL Likelihood =-3.03 Transmembrane 78 - 94 ( 75 - 94)

----- Final Results -----
bacterial membrane --- Certainty=0.6731(Affirmative) < succ>
45 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 9487> which encodes amino acid sequence <SEQ ID 9488> was also identified.

50 The protein has homology with the following sequences in the GENPEPT database:

```

>GP:CHA96187 GB:Z71552 AdcB protein [Streptococcus pneumoniae]
Identities = 197/263 (74%), Positives = 236/263 (88%)

```

Query: 13 LLDMLSYDFMQRAALLAVVAISIFAPILGIFLIIRRQSIMSDTLSHVSLAGVAGLVGLGIS 72
 +L +LSYDF+QRA LAV+A+S+F+P+LG FLIIRRQSIMSDTLSHVSL+GVA G+VLGIS
 Sbjet: 1 MLSSLGYDPIQRAFLAVIAMLSLPSVLGTFLIIRRQSIMSDTLSHVSLGVAFGVLGIS 60

Query: 73 PTWSTIFVVTLAAVVLEYLRVYKHYMEISTAILMGLAISLIMSKAINVGNVSLQY 132
 PT STI +V +AAV LKYLRTVYK +MSI TAILMS GLA+SLIMSK + ++SL+QY
 Sbjet: 61 PTVSTIAIVLIAAVFLEYLRVYKSPMEIGTAILMSTGLAVSLIMSKGSSSSMSLDQY 120

Query: 133 LFGSIITIGKEQVIALFVIALITFILLFIRPMYILTFDEDTAPVDGLFVRTMSILFNV 192
 LFGSI+TI +EQVI+LFVIA + IIL LF+RPMYILTFDEDTAPVDGLFVRTMSILFNV
 Sbjet: 121 LFGSIVTISEEQVLSLVIAAVALIILTFILFIRPMYILTFDEDTAPVDGLFVRTMSILFNM 180

Query: 193 VTGIAIALTIAPAGALLVSTINVLPASTIAMRLGRNFKTVIFLGMIGFVGMVAGIFLSY 252
 VTG+ALAL IPAGALLVSTINVLPASTIA+RLG+NFK+V+ L IGF+GMVAG++SY
 Sbjet: 181 VTGVAIALMIAPAGALLVSTINVLPASTIALRLGKNFKSVMLLSAIGFLGMVAGLYISY 240

Query: 253 WETPASATITMIFIGIFLLSLV 275
 ETPASA+IT+IF+ +F+L+SLV
 Sbjet: 241 AETPASASITIFVTVFILISLV 263

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 363> which encodes the amino acid sequence <SEQ ID 364>. Analysis of this protein sequence reveals the following:

Possible site: 18
 >>> Seems to have a cleavable N-term signal seq.

INTEGRAL	Likelihood	Transmembrane	135 - 151 (123 - 162)
INTEGRAL	Likelihood = -9.08	Transmembrane	68 - 84 (44 - 86)
INTEGRAL	Likelihood = -6.95	Transmembrane	20 - 36 (19 - 37)
INTEGRAL	Likelihood = -6.90	Transmembrane	251 - 267 (245 - 270)
INTEGRAL	Likelihood = -6.58	Transmembrane	221 - 237 (217 - 243)
INTEGRAL	Likelihood = -6.42	Transmembrane	91 - 107 (89 - 111)
INTEGRAL	Likelihood = -4.78	Transmembrane	176 - 192 (171 - 215)
INTEGRAL	Likelihood = -3.82	Transmembrane	45 - 61 (44 - 67)
INTEGRAL	Likelihood = -3.61	Transmembrane	196 - 212 (193 - 215)

----- Final Results -----
 bacterial membrane --- Certainty=0.6986 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP:CAA96187 GB:271552 AdcB protein [Streptococcus pneumoniae]
 Identities = 195/262 (74%), Positives = 239/262 (90%)

Query: 3 MLDILFYDFMQRAVMAVVAISIFAPILGIFLIIRRQSIMSDTLSHVSLAGVAGLVGLGIS 62
 ML +L YDF+QRA +AV+A+S+F+P+LG FLIIRRQSIMSDTLSHVSL+GVA G+VLGIS
 Sbjet: 1 MLSSLGYDPIQRAFLAVIAMLSLPSVLGTFLIIRRQSIMSDTLSHVSLGVAFGVLGIS 60

Query: 63 PTITTIIVVVIAAILLEYLRVYKHYMEISTAILMSGLALSILIMSKSHSSSSMSLEQY 122
 PT+TI +V++AA+ LEYLR VYK +MSI TAILMS GLA+SL+MSK SSSMSL+QY
 Sbjet: 61 PTVSTIAIVLIAAVFLEYLRVYKSPMEIGTAILMSTGLAVSLIMSKGSSSSMSLDQY 120

Query: 123 LFGSIITISGEQVIALFAIAAILITLVLFIRPMYILTFDEDTAPVDGLFVRTMSILFNI 182
 LFGSI+TIS EQV++LF IAA++LIL LF+RPMYILTFDEDTAPVDGLFVR MS+LNF+
 Sbjet: 121 LFGSIVTISEEQVLSLVIAAVALIILTFILFIRPMYILTFDEDTAPVDGLFVRTMSILFNM 180

Query: 183 VTGVAIALTIAPAGALLVSTINVLPASTIAMRLGKNFKTVILGIVIGFSGMLGIFLSYF 242
 VTGVAIAL IPAGALLVSTINVLPASTIA+RLGKNFK+V+LL IGF GM++G++ST+
 Sbjet: 181 VTGVAIALMIAPAGALLVSTINVLPASTIALRLGKNFKSVMLLSAIGFLGMVAGLYISY 240

Query: 243 FETPASATITMIFISIFLLSLV 264
 ETPASA+IT+IF++F+L+SL
 Sbjet: 241 AETPASASITIFVTVFILISLV 262

An alignment of the GAS and GBS proteins is shown below:

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Identities = 223/270 (82%), Positives = 252/270 (92%)

Query: 12 MLLDMLSYDFMQRALLAVVAISIFAPILGIFILRRQSLMSDITLSHVSLAGVALGVVLGI 71
 ++LD+L YDFMQR++AVVAISIFAPILGIFILRRQSLMSDITLSHVSLAGVALGVVLGI
 5 Sbjct: 2 VMLDLILYDFMQRVAVVAISIFAPILGIFILRRQSLMSDITLSHVSLAGVALGVVLGI 61

Query: 72 SPTWSTIFVWTLAAVLELYLTVKHYMEISTAILMSGLAISLIVMSKAHNVGNVLSQ 131
 SPT +TI VV LAA++LEYLR VYKHYMEISTAILMS+GLA+SLI+MSK+H+ ++SLSQ
 10 Sbjct: 62 SPTTTITIVVLAAILLEYLRVYKHYMEISTAILMSGLAISLIMSKSSSSMSLBQ 121

Query: 132 YLFGSIITIGKEQVIALFVIAIITFILITLIRPMYILITDREDTAFVDGLPVRIMSVLEN 191
 YLFGSIITI EQV+ALF IA I ILT+LIRPMYILITDREDTAFVDGLPVR MS+LFN
 Sbjct: 122 YLFGSIITISMEQVVALFAIAAILILTLVLRPMYILITDREDTAFVDGLPVRIMSVLEN 181

Query: 192 VVTGIAIALITIPAGALLVSTIMVLPAIAMRLGRNFKTVIFGLMIGFQVMVAGIFLSY 251
 +VTG+AIALITIPAGALLVSTIMVLPAIAMRLG+NFKTVI LG++IGF GM++GIFLSY
 15 Sbjct: 182 IVTGVAIALITIPAGALLVSTIMVLPAIAMRLGRNFKTVILLGIVIGFSGMLSGIFLSY 241

Query: 252 YWETPASATTIMIFIGIFLVLVGLLRKR 281
 ++ETPASATTIMIFI IFLVSL G+L+KR
 20 Sbjct: 242 FFETPASATTIMIFISIFLVLVGLMLKKR 271

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

25 Example 110

A DNA sequence (GBSx0113) was identified in *S.agalactiae* <SEQ ID 365> which encodes the amino acid sequence <SEQ ID 366>. This protein is predicted to be streptodornase. Analysis of this protein sequence reveals the following:

Possible site: 59
 30 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.2601(Affirmative) < succ>
 35 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAA59264 GB:X84793 streptodornase [Streptococcus pyogenes]
 40 Identities = 58/167 (34%), Positives = 85/167 (50%), Gaps = 30/167 (17%)

Query: 2 TPIYEGNMLVPSRVELQYVGIDKQKLEIKLGGKQVDEYGVTTVLENTSPLAKIDY 61
 TP+Y+G+ L+P V + + D +DE TV + N IDY
 45 Sbjct: 245 TPVYQSGELLPRAVLVSAALSSDGF-----IDE---TVRVFNNAVGFNTDY 286

Query: 62 KTGMLIKEDGKQAEEDGPDNSADENGAIR-SASDIERNNTINTSESOTNVAPQIRIV 120
 + G L+ E P + D E +E + IE+ +T+T + D N++ Q + V
 50 Sbjct: 287 QNGKGLTSS-----PVITETNVSRNVEDNITETIDEVDITDLKKIDENISLQ-KTV 336

Query: 121 YVANKGRSNTYWYSLENI-KNMTANIYQMTQRQALNKHHSITTEA 166
 YV+ G SN YWYS EN+ KN N +V+M+EQ AL + KHS EA
 55 Sbjct: 337 YVASSGLSNVYYSKRNMPKNVNLQKVMESRQTALRGKHSAQEA 383

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 367> which encodes the amino acid sequence <SEQ ID 368>. Analysis of this protein sequence reveals the following:

Possible site: 31

>>> Seems to have a cleavable N-term signal seq.

-178-

----- Final Results -----

bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

5

An alignment of the GAS and GBS proteins is shown below:

Identities = 51/90 (56%), Positives = 66/90 (72%), Gaps = 4/90 (4%)

Query: 1 MTPIIVEGNLVESEVLEQVIGIDKQKLEIKLGGKKQVDEYGVITVLENTSPFLAKID 60
 +TF+Y N LVP +V LQVVID+ G LL+IKLG KR VD +QVT+VTL+N SFLLA+D
 Sbjct: 182 VTFVYHKGSLVFRQVVLQVIGIDENGDLQIKLGSEKSEVNFQVTSVTLNVSFLAELD 241

Query: 61 YKTGMILIKEDKQKQREGEFNSDADENRAA 90
 Y+TGM++ D Q E ED N + +E E A
 Sbjct: 242 YQTGMML--DSTQNE--EDSNLETRFSEA 267

15

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 111

20 A DNA sequence (GBSx0114) was identified in *S.agalactiae* <SEQ ID 369> which encodes the amino acid sequence <SEQ ID 370>. This protein is predicted to be tyrosyl-tRNA synthetase (tyrS-1). Analysis of this protein sequence reveals the following:

Possible site: 60

25

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3618 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

30

The protein has homology with the following sequences in the GENPEPT database:

>GP:AA00303 GB:AF008220 tyrosine tRNA synthetase [Bacillus subtilis]

Identities = 234/420 (55%), Positives = 311/420 (73%), Gaps = 2/420 (0%)

35

Query: 2 NIFDELKERGLVFTQTEDALRKALEEGSVSYTGYDPTADSLHGLHVAITLSRRLQLA 61
 N+ ++L RGL+ Q TDE+ L K L E + Y+G+DPTADSLH+GHL+ ILT RR QLA
 Sbjct: 3 NLLEDLSFRGLIQMTDEEGLNKQLEEKIRLYSGCPDPTADSLHGLHLPILTLLRFPQLA 62

40

Query: 62 GHKPYALVGGATGLIGDPSFKDVERSLQTKKIVSVWGNKIRQQLNSFLEFQTDGNKAVLV 121
 GH P ALVGGATGLIGDPS K ER+L T V W KI+ QLS FL+FE +N AV+
 Sbjct: 63 GHHPIALVGGATGLIGDPSGKKAERTLTNTADIVSEWSQKIKNQLSRFLDFEAENPAVIA 122

45

Query: 122 NNYDNFNSISIFDLRDVGKQVPTVNMVMSKESVKRIETGISITFEAYIQMGQYDFVELN 181
 NN+DN ++ IDFLRDVGK F +NVM++K++V RIE+GISITFE+Y I+Q YDF L
 Sbjct: 123 NNFCWIGKGNVIDFLRDVGKNFGINVMALKDPTVSSRIEISGITFEYSYMLQSYDFNLNLY 182

50

Query: 182 KNYNVTLQIGSSDQWGNMAGTGLIRK--KSNQSVHVMVTPLITDSTGKKGKSEGNVAVN 239
 ++ N LQIGSSDQWGN+TAG ELIR+ + + +T+PL+T+ G KPGK+EG A+W
 Sbjct: 183 RDNCKLQIGSSDQWGNMAGTGLIRKSESGAKAFLGTIPLVTKADGTGKGTGGGAIN 242

55

Query: 240 LDADKTSFYEMQYQWLNMDADAVRFLKIPTPLSLKEIEDIRIQFEAPHQLAQKTLAR 299
 LD +KTSFYE YQW+N D D V++LK PFLS +EIE + E AP +R AOK LA
 Sbjct: 243 LDKEKTSFYEFYQFWLNTDTRDVVYLYKYFTPLSKREIRAYARCTEATPGRFAQKRLAE 302

Query: 300 EVVTLVHGEKAYKEAVNITQFLFAGNIKLSVKRLKQGLRGVPHYVCTENMNIIDILV 359
 EV +LVHG +A ++A+NI++ LF+GNIK LS +++K G + VP+ V + L+++D+LV
 Sbjct: 303 EVTSLVHGREALEGAINISQALFSGNIKLSKADQVVKVGFQVPSMEVDSTQELSLVDVLV 362

60

Query: 360 TSGVNSKRQAREDSNGAIYINGDRIQLEVTYSNDKLENETTVIRGKKYCVLNFK 419

-179-

S + SERQARE+ NGA+YING+R ++ YT+S D++EN+ TV+RRGKKKYF++ +K
 Sbjct: 363 QSKLSPSKRQAREDIQNGAVYINGRQTEINITYLSGRDRIENQFTVLRGKKKYFLVITYK 422

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 371> which encodes the amino acid sequence <SEQ ID 372>. Analysis of this protein sequence reveals the following:

Possible site: 37

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2340 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 344/418 (82%), Positives = 377/418 (89%)

Query: 1 MNIPDELKERGLVFQTTDEDALRKALEDGGSVSYTYGYDPTADSLHLGHLVAILTSRRLQL 60
 MNIP+ELK RGLVFQTTDE AL KAL EG VSYTYGYDPTADSLHLGHLVAILTSRRLQL
 Sbjct: 1 MNIPDELKARGLVFQTTDEQALVKALTEGQVSYTYGYDPTADSLHLGHLVAILTSRRLQL 60

Query: 61 AGHKPYALWGGATGLIGDPSFKDVERSLQTKTKTVVSGNKRQGLSNFLEPETGDNKAVL 120
 AGHKPYALWGGATGLIGDPSFKD ERSLSQTK+TV+ W +KI+GQLS FL+FE GDNKA L
 Sbjct: 61 AGHKPYALWGGATGLIGDPSFKDAERSLQTKETVLEWSDKIKGQLSTFLDFENGDNKAL 120

Query: 121 VNNYDWFNSISFIDFLRDVGKYFTVNNYMSKESVKKRIETGISYTEFAYQIMQGYDFYEL 180
 VNNYDWFNS ISFIDFLRDVGKYFTVNNYMSK+SVKKRIETGISYTEFAYQIMQGYDFYEL
 Sbjct: 121 VNNYDWFNSISFIDFLRDVGKYFTVNNYMSKESVKKRIETGISYTEFAYQIMQGYDFYEL 180

Query: 181 NKNYNVTLQIGGSDQMGWNTAGTELLRRKSGVSHVMTVPLITDSTGKKPGKSGNAVWL 240
 N +NVTLQIGGSDQMGWNTAGTELLR+K++ HVMVTPLITDSTGKKPGKSGNAVWL
 Sbjct: 181 NDKGNVTLQIGGSDQMGWNTAGTELLRRKKADKTHVMTVPLITDSTGKKPGKSGNAVWL 240

Query: 241 DADKTSPYRMYPQFWLAVMDADAVRFLKIPTFLSLKEIDIRIQFRRAPHRQAQKTLARE 300
 DADKTSPYRMYPQFWLAVMD DAVRFLKIPTFLSL EI +I QF A H+RLAQKTLARE
 Sbjct: 241 DADKTSPYRMYPQFWLAVMDDDAVRFLKIPTFLSLDEIAEITQFNAARHERLAQKTLARE 300

Query: 301 VVTLVHGEEKAYKEAVNITEQLPAGNINKLSVKELKQQLGAGVPNHYVQSDNINMIDLLVT 360
 VVTLVHGE+AYK+A+NITEQLPAGNIK LS ELKQQL VPNYHVQ+ DN NI+++LV
 Sbjct: 301 VVTLVHGEEAYKQALNITEQLPAGNINKLSANELKQQLGSLVPNHYVQSDININMIEVLVA 360

Query: 361 SGVNSKRRQAREDSNGAIYINGDRIQDLEYTISENDKLENEITVIRRGKKKYFLVNL 418
 + + SKRQAREDV NGAIFYINGDR+QDL+Y +S +D++++TVIRRGKKKY VL+
 Sbjct: 361 AKISPSKRRQAREDVNGAIYINGDRVQDLQYQLSNDKIDQLQTVIRRGKKKYVLVITY 418

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 112

A DNA sequence (GBSx0115) was identified in *S.galactiae* <SEQ ID 373> which encodes the amino acid sequence <SEQ ID 374>. Analysis of this protein sequence reveals the following:

Possible site: 53

>>> Seems to have no N-terminal signal sequence

INTEGRAL Likelihood = -12.21 Transmembrane 36 - 52 (23 - 59)

----- Final Results -----

bacterial membrane --- Certainty=0.5883 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

-180-

The protein has homology with the following sequences in the GENPEPT database:

>GF:AA04736 GB:AF101781 penicillin-binding protein 1b
[Streptococcus pneumoniae]
Identities = 445/769 (57%), Positives = 581/769 (74%), Gaps = 9/769 (1%)

Query: 3 KKNKLNLSKIGDYTF---LEFGSIFLRI---VKLLSDFIYVILLFVMLGVLAVGYL 55
K K K G T L+ +IF I +K L+ ++V+ L M6 G+A+GY
Sbjct: 21 KKNKSARPGKSGSSTKSKTSLDKSAIPAILLSIKALPNLLFVLGFLGQMLGAGLAVGY 60

Query: 56 ASQVDSVKVPSKNSLVTQVNTLTVRSRLITYSDKQISEIATDLQRTFVKADAIISNIIKA 115
+ D V+VP LV QV ++ +S +TYS D + I + DL RT ++ + IS+N+KKA
Sbjct: 81 VALPDKVRVPQTELVNQGKDISSISEITTSYDGTVIASIESDLKRTSISSEIQISENLKKA 140

Query: 116 IIAITEDENFNDHKGVVPKAVLRRAAGSVLGFSGESGGSTLTQQLLKQQLIGDDPSFKRKS 175
IIATEDE+P +HKGVPKAV+RA G +G G SSGGSTLTQQL+KQ++GD P+ R+
Sbjct: 141 IIAATEDEHFKHKGVPKAVIRATLGFVGLSGSGGGSTLTQQLIKQVGVGDATLARKA 200

Query: 176 KEIIYALALERYMDKSDLSIDYLVNVPFORNNKQNIAGIEBMAQSIGVSAKDLTIPCA 235
EI+ ALALER M+KD IL+ YLVN+PFGNNKQNIAG ++A+GIFGV A LT+PCA
Sbjct: 201 AEIVDALALERAMKDEILITLVNVA+PGRNNKQNIAGARQAAGSIGVDSQLTVPQA 260

Query: 236 AFLAGLPQSPFIVSYSPYTADAQLKSDKDLSPGIFKQKNVLYMYRTRALTQKDEYSKYDYD 295
AFLAGLPQSPI YSPY +LKSD+DL G+R K VLY+MYRT AL+KDEY KYDYD
Sbjct: 261 AFLAGLPQSPITISPYENTGELKSDLEDLEGLERAKAVLYSMYRTALSKDEYSKYDYD 320

Query: 296 IKQDFIKPAVATTNHEDVLYSALSEAQKVMYNYLKKDNVSEEDLNDIDETRATYRHRAI 355
+K+DP+ T DYLV++ L+EAQ+ MY+YL +DNVS +LQN+ T+ YR A
Sbjct: 321 LKQDFLPSTGTGTSIDYLVFTTLAEAQERMYDYLQKDNVSAKELQETATQFYRDLAA 380

Query: 356 EEIQQGGYTIKTTINKSVYQAMQDRAAQYGLLDDGTGKQVMGNVLTNNSGAILFGIG 415
+EI+ GGY I TTI++ ++ AMQ A A YG LLDDGTG+V++GNVL DN +GAI+GF+GG
Sbjct: 381 KEIBNGGYKITTTIDQKIHSAQSAVADYGYLLDDGTGRVEGNVLMNDQNGAILGFVG 440

Query: 416 RNYSENQNNHAFDTARSQSSIKPILPYGIAIDQGMGSGSVLSNPTYPTSSGRKIMHAD 475
RNY ENQNNHAFDT RSP S+ KP+L YGIAIDQ++GS ++LSNPTY ++G IM+A+
Sbjct: 441 RNYSENQNNHAFDTRSPASTTKPLLAYGIAIDQGMGSETILSNPTNFANGNPIMYAN 500

Query: 476 REGTAMVNLQESLDISWNIAPWYTYMLRDGVDVKNYMEKLDYPIENFQIESLPGGGI 535
+GT M+ L E+L+ SWNI PA+WTY+MLR+ GVDVK YME+ Y I +GIESLP+GGGI
Sbjct: 501 SKGTGMNTLGEALNYSWNI PAWYTYMLREKGVVDVKNYMERMGYEIPEYQIESLPMGGI 560

Query: 536 DTSVAQQTINLYQMIANGGVYHQYKYMIESIEDSGNKVLYNHSEKPVRVFSKATATILQQL 595
+ +VAQ TN YQ +AN GVYH++I IE ++G+V+Y ++ KPV+V+SKATATI+Q LL
Sbjct: 561 EVTVAQHNGYQTLANNGVYHQKHVLSKIEBADGRVVVEYQDKPQVVSFKATATILQQL 620

Query: 596 HGPINSKTKITTFKRLQGLNSGLAGVDMWIGKTGTNTSDVMMLSTPKVTLGSMAGHDN 655
++S TITTFK+ L LN LA DWIGKTGTN ++NMLSTP++TLGSG GHD+
Sbjct: 621 REVLSRVTITTFKSNLTSLMPTIANADWIGKTGTNTQDNMMLSTPRLTLGSLMGHD 680

Query: 656 NASLAKLTGYNNNANYMAHLVNAINNADGRTFGKSERFLDDSVIRAKVLKSTGLQPGVV 715
N SL++ GY+NN+NYMAHLVNAI A + +G +ERF LD SV+K++VLKSTG +PG V
Sbjct: 681 NMSLSRAGYSNNANYMAHLVNAIQASPSIWG-NERFALDPSVVKSEVLKSTGQKPGKV 739

Query: 716 TVNGRRLTVGSESTTSYWA-KNGPGTMYRPAIGQTDSDYKAWSTLGG 763
+V G+ + V G+ TSYWA K+G +YRPAIGG++DYQ AWS++ G
Sbjct: 740 SVEGKEVTVGSTVTSYVANKSGAPATSYRPAIGGSDADYQNAWSIVG 788

A related DNA sequence was identified in *Spyogenes* <SEQ ID 375> which encodes the amino acid sequence <SEQ ID 376>. Analysis of this protein sequence reveals the following:

Possible site: 57

>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood = -4.83 Transmembrane 39 - 55 (32 - 60)

----- Final Results -----

-181-

bacterial membrane --- Certainty=0.2932(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

- 5 The protein has homology with the following sequences in the databases:

>GP:AAF04736 GB:AF101781 penicillin-binding protein 1b
 [Streptococcus pneumoniae]
 Identities = 438/739 (59%), Positives = 580/739 (78%), Gaps = 2/739 (0%)

10 Query: 27 FVLRRTLRLLSNFFYIVIEFLPGMAGPQMAPGYLASIQSVKPSKESLVKQVRSLLTMIQ 86
 P +L +++ L N +++ FL GM+G G+A GY + + V+VP E LV QV++ ++ IS+
 Sbjct: 48 PAILLGLKALPNLLPVGLGFLGMLGAGIALCYGVALFDKVRVQTERLVNQVDDISSISE 107

15 Query: 87 MYGINSGLISTLDITLLRTFVNDIAISENKKIAVSTEDEHFQEHKGIVPKAVFRATLAS 146
 + YSD ++I++++DILLRT ++++ ISRN+KAI++TEDEHP+ERHKG+VPKAV RATL
 Sbjct: 108 ITYSDGTVIASIESDILLRTSISSEQISENKKAIATEDEHFQEHKGIVPKAVIRATLCK 167

Query: 147 VLGGFASGGSTLTQQLVKQVGLGDDPTFRKSKSEIVTALALERYMSKDNILCDYLVSRP 206
 +G G +GGGGSTLTQQL+KQV+GD PT RK+ EIV ALALER M+KD IL YLV+P
 Sbjct: 168 FVGLGSSGGSTLTQQLIKQVVGDAFLTKARKAIVDALALERAMNDEILTYLVNAP 227

20 Query: 207 FGRNKGQNTIAGVEAARGIGVSAKDLTPVQAAPLAGLPQSPITVYSPYLSTOQLKSEKD 266
 FGRNKGQNTIAG +AA GIGV A LTPVQAAPLAGLPQSPITV YSPY +TG+LKS++D
 Sbjct: 228 FGRNKGQNTIAGARQAABGIGVSDASQLTPVQAAPLAGLPQSPITVYSPYENTGELKSED 287

25 Query: 267 MAYGIRKQNVLPFRYRTGVLSKEYEDYKAYPIQDKIQPGSAIVNNHDLTYTVLADA 326
 + G+R + VL++MYRTG LSK EY YK Y +++DF+ G+ + DYLY+T LA+A
 Sbjct: 288 LEIGLRAKAVLYSNMYRTGALSKEYEDYKAYPIQDKIQPGSAIVNNHDLTYTVLADA 347

30 Query: 327 KIMAYSYLLKRDKVSRLDNDETKAAYBERALTELQOQGYTITTTINKPIYNAMOTRAA 386
 ++ MY YL +RD VS++LKN+ T+ Y + A E++ GGY ITTII+ I++AMQ+A A
 Sbjct: 348 QERMYDYLAQRDNVSAKELKNEATQKPYRLAAKEIENGQYKITTITDQKIHBAQSAVA 407

35 Query: 387 QFGGLDDGTGTGVGNVLTNDATGAVLGFVGRDVALNQNNAHNPVTSRSGSSKPIIA 446
 +G LDDGTG V++GNVL LN TGA+LGFVGR++Y NQNNHAF+T RSP S+ KP++A
 Sbjct: 408 DYGVLDDGTGRVGNVLTNDATGAILGFVGRNRYQENNNHAFDTRKSPASTTKPLIA 467

40 Query: 447 YGALDQGLMGSGASVLSNYPTTYSQQKIMHADSEGTAMMPLQALNTSNIPAFWTKL 506
 YG AIDQGLMGS ++LSNYPT +++G IM+A+S+GT MM L EALN SNNIPA+WT ++
 Sbjct: 468 YGALDQGLMGSETLSNYPTNFAHGNIPTVANGSGTGMMLGSEALNYSNNIPATWITYRM 527

45 Query: 507 LREKGVVDVENVMTKMGYKIADYSIESLPGGGIEVSAQQTNAVQMLNNGLYQKQYVD 566
 LRE GVDV+ YN RMGY+I +Y IESLP+GGGIEV+VAQ TN YQ L+NG+Y +++++
 Sbjct: 528 LREKGVVDKGYMEKNGYIEPEYQIESLPGGGIEVTVAQHTNGYQLANNGVYHGVKIVIS 587

Query: 567 KITASDGTVVYKHNKPIRIPSAATATILQELRGPIITSGATTTFKNLAAINPWLANAD 626
 KI A+DG VVY++++KP++++S ATATI+Q LIR ++S TTTFFK+ L ++NP LNDAD
 Sbjct: 588 KIEAADGRVYVEYQDKPVQYSKATATIMQLREVLGSRVTTTFKNLSINPWLANAD 647

50 Query: 627 WIGKIGITENYTDVNLVSTPKVTLGQWAGHEDNISLAPLTGYNNNENYLAAYANAINQA 686
 WIGKIGIT ++WL+LSTP++TLGQW GHEDN SL+ GY+NNENY+A+L NAI QA
 Sbjct: 648 WIGKIGITNDQENGLMLSTPRLTGLGWIGHEDNHELRRAGYNNENYMHVNLVAINQA 707

55 Query: 687 DPNVIGVGQRFLNDPGLVKANVLKSTGLPGQTVNVNGHTFSGVGGEMTSLMSQK-GPGAM 745
 P++ G +RF LDP V+K+ VLKSTG +PG V+V G V G TS W+ K G A
 Sbjct: 708 SPSTIWG-NERFALDPSVVKSEVLKSTGQKPKGVSVEGKEVEVTGTSVTSYWNKSGAPAT 766

Query: 746 TYRFAIGGTDADYQKAMWN 764
 +YRFAIGG+DADYQ AM +
 Sbjct: 767 SYRFAIGGSDADYQAMBS 785

An alignment of the GAS and GBS proteins is shown below:

Identities = 531/760 (69%), Positives = 639/760 (83%), Gaps = 3/760 (0%)

65 Query: 6 KKLNSSKLGDYTFLEPGSIFLRIVKLLSDFTYVILLFVLMGLAVGLAVGYSQVDSVKVP 65

K+++ +LG L+ G + LR ++LLS+F Y++I LF M+G G+A GYLASQ++SVKVP
 Sbjct: 13 KRISHQRQLG---LLDLGVLLRTLRLLSNFFYIVIFLPGMGMFGNAPGLASQIESVKVP 69

5 Query: 66 SKNSLVTQVNTLRVSRLTYSKDSQISEIATDLQRTFPAKDAISDNLKKAIIATRDENFN 125
 SK SLV QV +LT +s++ YSD S IS + TDL RFPVA DAIS+NLKKAII+TEDE+F
 Sbjct: 70 SKESLVKQVESLTMISQBNYSNSLISLTLDTLLELTFVANDAISENIKKAIVSTDESHFQ 129

10 Query: 126 DHKGVVPEKAVLRAAASVLPFGSSGGSTLTQQLLQQLIGDDPSVKRSKEIYALALE 185
 +HRG+VPEKAV RA SVLPFGSE+GGSTLTQQL+KQ+LGDDP+FKRSKEI+YALALE
 Sbjct: 130 RHKGIVPEKAVFRATLAVLPFGKASGGSTLTQQLNQQLVIGDDPTFKRSKEIYALALE 189

15 Query: 186 RYMEDKDSILSDYLVNSPPGRNNKNGQNIAGIEKKAAGIPGVSAKDLTVPQAFLAGLPQSP 245
 RYM KD+IL DYLNVSPFGRNNKNGQNIAG+EEAA+GIPGVSAKDLT+PQAFLAGLPQSP
 Sbjct: 190 RYMSEKNTLCDYLVNSPPGRNNKNGQNIAGVEKAARGIPGVSAKDLTVPQAFLAGLPQSP 249

20 Query: 246 IVISPYTADACLKSKDKLSPGIKQKQNVLYNMTRFALCKDEYKSKYDQIDKDFIKPAV 305
 IVISFY + QLSK+KD++GIKRQ+NVL+NMTR L+K EY+ YK Y I+KDFI+F
 Sbjct: 250 IVISPYSLTQGLSKRDMVGIKQKQNVLFNMTRTGVLSKKEYEDYKAYPIQDFIQPGS 309

25 Query: 306 ATTNRHDYLYYSALSEAQKVMYNYLIKKDNVSEHLDKNDETSTRATRYHERAIEIQGGYTI 365
 A N+HDYLYY+ L+A+A+K MY+YLIK+D VS DLKDET+A Y RA+ E+QGGYTI
 Sbjct: 310 AIVNRHDYLYYTLADAKKAMYSTYLIKRDKVSRLKNDETKAAYTEERALTELQGGYTI 369

30 Query: 366 KTTINKSVYQAAQDAQAQYGLLDDGTGTVQNGVNLTDNNSGAIIGFIGNYSNQNH 425
 TTINK +Y ANQ AAAC+QGLLDDGTG VQNGVNLTDN++GA++GGR+Y+ NQNH
 Sbjct: 370 TTTINKPIYNAAQDAQAQYGLLDDGTGTVQNGVNLTDNATGAVLFGVGRDYALNQNH 429

35 Query: 426 AFDTARSPGSSIKPILFYGLIALDQMLSGSGLVSNYPITYSSGKINHADEBGTAMVNLQ 485
 AF+T RSPGSSIKPI+ YG ALDQ++GS SVLSNYPITYSSG+KINHAD EGTAM+ LQ
 Sbjct: 430 AFTVRSFGSSIKPIIAYGALDQGLMGSASVLNYPITYSSGQKINHADEBGTAMNPLQ 489

40 Query: 486 ESLDISWNIAPFWTYMLRDRGSDVINYMEKLDYIFNFIQIESLPLGGGIDTSVAQQTNL 545
 E+L+ SWNIAPFWT K+LR++GVDV+NYM K+ Y I ++ IESLPLGGGI+ SVAQQTN
 Sbjct: 490 EALNTSWNIAPFWTQKLREKGVDFVNYMTNGYKTADYSIESLPLGGGIEVSVAAQQTNA 549

45 Query: 546 YQMIANGGVYHRYMIESIEDNGKVIYNHESKIVRVFVSATATILQQLLHGFISNGKTT 605
 YQM++N G+Y KQY+++ I S+G V+Y HE+KP+R+FS ATATILQ+LL GP SG TT
 Sbjct: 550 YQMLNNGLYQKQYIVDKITASDGTVVYKHENKPIRIFSAATATILQELLRGPITSGATT 609

50 Query: 606 TFIKRLQGLNSGLAGVDWIGKTGTINSTDVWMLSTPKVTILGGWAGHDNNSLAKLTGY 665
 TFIKRL +N LA DWIGKTGT + +DVWL+LSTPKVTILGGWAGHD+N SLA LAGY
 Sbjct: 610 TFIKRLAALPNLANADWIGKTGTETRYTDVWLVLSTPKVTILGGWAGHDNTSLAFLTGY 669

Query: 666 MNAYMYAHLVANAINNADGMTFGKSERFLDSDVIKAVKLTSGTLGPQVTVNGRRITVG 725
 MN+NY+A+L NAIN AD N G +RF LD VIKAVLKSTGLQPV VNG +VG
 Sbjct: 670 MNSNYLAVLANAINDADPMVIGVGRFNLDPGVKANVLKSTGLQPVTVNGRHITFVG 729

Query: 726 GESTTSYWAINGPGMTMYRFAIGGTDSDYQKANSLLGGRK 765
 GE TTS W++ GGR MTYRFAIGGTD+DYQKAN G ++
 Sbjct: 730 GEMTTSLSNSQKPGAMTYRFAIGGTDADYQKANGNRFPRK 769

SEQ ID 374 (GBS64d) was expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 120 (lane 2-4; MW 107kDa). It was also expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 120 (lane 5-7; MW 82kDa) and in Figure 179 (lane 2; MW 82kDa).

GBS64d-His was purified as shown in Figure 231, lane 7-8.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 113

A DNA sequence (GBSx0116) was identified in *S.agalactiae* <SEQ ID 377> which encodes the amino acid sequence <SEQ ID 378>. This protein is predicted to be DNA-dependent RNA polymerase subunit beta (rpoB). Analysis of this protein sequence reveals the following:

```

5   Possible site: 61
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
10  bacterial cytoplasm --- Certainty=0.3505(Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
    bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

15  >GP:CRB56706 GB:Y16468 DNA-dependent RNA polymerase subunit beta
    [Listeria monocytogenes]
    Identities = 814/1173 (69%), Positives = 978/1173 (82%), Gaps = 17/1173 (1%)

    Query: 2   AGHEVQYGHRTKRSFSPRIKEVLDLNLIEIQDTSPOQFLDAGLKEVFDVLPISFDT 61
    +GH+Y+YGHRTKRSF+RI EVL+LNLIEIQT S+Q FLD GL+E+P D+ PI +F
20  Sbjct: 5   SGHDVKYGRHRTKRSFARISEVLELNLIEIQIASYQWFLDGLREMPRIDISPFAGN 64

    Query: 62   MDLEFVGVELKEPKYITLLEARIHDASYSAPIFVTFRLNKEITGEIKTQEVFPGDFPMTTE 121
    +LEF+ Y+L EPKY++EE++ DA+Y+AP+ V RL+NKEIGE+K QEVF GDFP+TE
25  Sbjct: 65   LSLFIDYDLGEPKYSVESSEKRDANYAAPLRVKLRINKETGEVKDQEVFPGDFPLATE 124

    Query: 122  MGTFTINGGERIIVSQLVRSPGVFNKDKNKGKGVGSGTVIPNRGAMLETDADJLAY 181
    MGTFTING ER+IVSQLVRSPGVFN K+DNKKG G+GSTVIPNRGAMLETDADJLAY
30  Sbjct: 125  MGTFTINGAERVIVSQLVRSPGVFNKGLDNKKGKGVGSGTVIPNRGAMLEYETDAKOVVH 184

    Query: 182  TRIDTRKIPPTTLVRLALGSGDDEIVDIPGDSIAVNTIEKDIDHNKPSDSSTDEALKEI 241
    RIDTRK+P T L+RALSF D EIT+D GD++ +ENT+EKD N ++AL EI
35  Sbjct: 185  VRIDTRKLPVTVLLRALGSGSDQEIIDILGNDVLENTLEKDNNT-----AEKALLEI 239

    Query: 242  YERLRPGEPKTDSSRLIARFPDPFRDYDLAAGVGRKINKKLNKTRLNQTAENLVD 301
    YERLRPGEP T D++RSLIV+RFPDP+RYDLA+VGRYKINKKL+LK RL NQT+AR LVD
40  Sbjct: 240  YERLRPGEPPTVDNARSILVSRFPDPKRYDLASVGRYKINKKLNKTRLNQTAENLVD 299

    Query: 302  GETGEILLVEAGTVMTRDVIDSIAEHIDGLNKFVYTPNDYAVVTEPVILQKFVVAAPTD 361
    ETGEI+ G ++R +D I +++ + P D V+ +V ++K +P D
45  Sbjct: 300  PETGETIASKGDIILRRNLQIILNLENGVGFRTLRPTD-GVMEDSVIAVQS+KIYVNPDE 356

    Query: 362  DRVVTIVGNSNPEDKVEALTPADILARMGYFINLBSIGIKVDIDHGNRRIRAVGELL 421
    ++ + I+GN+ E+ V+ +TP+DI++ +SYF NL G+G DDIDHGNRR+R+VGEILL
50  Sbjct: 359  EKEINILIGNAVIERNVKHITPSDIISISYFNLLHGVQVDIDIDHGNRRIRAVGELLQ 416

    Query: 422  NQFRIGLRMRERNVRERMSVQONEVLTPQCIINIRPVTAAKEFPGSSQLSQFMDGNPL 481
    NQFRIGL+RMR VREEMS+QD +TQCG+INIRPV A++KEFPGSSQLSQFMDQ NPL
55  Sbjct: 419  NQFRIGLRMRERNVRERMSVQONEVLTPQCIINIRPVTAAKEFPGSSQLSQFMDQINPL 478

    Query: 482  SELSHKRRLSALGPGGLTRBAGYEVVRDVHYTHYGRMCPIETPESNIGLINSLSFGHL 541
    EL+HKRRLSALGPGGLTR+RAGYEVVRDVHY+HYGRMCPIETPESNIGLINSLSFGHL
60  Sbjct: 479  GELTHKRRLSALGPGGLTRBAGYEVVRDVHYTHYGRMCPIETPESNIGLINSLSFGAKV 538

    Query: 542  NKYGFIQTPYRKVDSTGAVINELVLTAEDEFTVAQANSKLNEGDTFAEELVGRH 601
    NK+GFI+TPYR+VD T VT++I +LTAEED + VAQANSKL+E GTF EE VM R +
65  Sbjct: 539  NKYGFIEPTPYRKVDSTGAVINELVLTAEDEFTVAQANSKLNEGDTFAEELVGRH 598

    Query: 602  GNNCEFPSSIVDFVDVS PKQVAVATACIPFLENDGSRNALGANMQRQAVPLDIPKAPY 661
    N +D++DVS PKQVV+VATACIPFLENDGSRNALGANMQRQAVPLDIPKAPY
70  Sbjct: 599  SENLAVEKERIDYMDVS PKQVVSVATACIPFLENDGSRNALGANMQRQAVPLDIPKAPY 658

    Query: 662  VGTGMEXQAANDSGAAVTAKHGIVHFAREIIVRRVSLVDGKEVVGIDKYTYLAKFVR 713
    VGTQME+ +A DSGAAV AKHDG V +A ++ VRR G+D Y ++K+P R
75  Sbjct: 659  VGTGMEXVAKDSGAAVTAKHGIVHFAREIIVRRVSLVDGKEVVGIDKYTYLAKFVR 718

```

Query: 714 SNSGTAYNQRTLVKVGDLVEKGFADGSPMENGEMALGNFVAYMTWESYNFEDAVIM 773
 SN GT YNQR V GD V KG+ + +GPM+ +GE+ALG+N +VA+MTW+SYN+EDA+IM
 5 Sbjet: 719 SNQSTCYNQRPVABGIRVVKGRILGNQSPMDSGKLAGRNVLVAPMTWQYNYEDAIM 778

Query: 774 SERLVKEDVYTSVHLKEFSESTRUTKLGPEETREIPNVGEDSLRDLDEMSILRIGAEVK 833
 SERLVK+DVYTS+H+EEFSE RDTKLGPEE+TR+IPNVGED+LRDLDE GIIR+GAEVK
 Sbjet: 779 SERLVKEDVYTSIHIEFSEARDTKLGPEEMTRDIPNVGEDALRDLDESGILRIGAEVK 838

Query: 834 EGDILVGVKVTPEKGEKDLSEAFERLLHAIFGDKSREVRDTSLRVPHGGDGVVRDKPIFRAN 893
 + D+LVGVKTPKG +L+AEERLLHAIFG+K+REVRDTSLRVPHOG G+V DVKPIFR
 10 Sbjet: 839 DNDLLVGVKVTPEKVTETAEERLLHAIFGKREVRDTSLRVPHGGGIVLVKPIFREAN 898

Query: 894 GDELQSGVNMILVRVYIAQGRKIKVGDGMAGRHGKGVSRIVPVEDMPYLPDGTVPVDIML 953
 GDEL GVN LVRVYI QGRKI GDMAGRHGKGV+SRI+P EDMF++PDGTVPVDIML
 15 Sbjet: 899 GDELPFGVNLVRVYIVQGRKIRHGDMAGRHGKGVSRILPEEDMPFMDGTVPVDIML 958

Query: 954 NPLGVFSRMNIGQVMELHGLNMAARNLGIHATVPFVDGASSEDMLWTVQEAQMDSDAKTVL 1013
 NPLGVFSRMNIGQV+ELHGLNMAAR LGIH+ATVPFDGA+ ED+W TV+EAM DAKT+L
 20 Sbjet: 959 NPLGVFSRMNIGCVLEHGLNMAARLGIHVATVPFDGANEDVNSTVEEAGVARDAKTIL 1018

Query: 1014 YDGRKGEPFNNRVSQVGYMYMKLHHMVDKLGHARSVGPYSLVTTQPLGGKAQGGQRPGE 1073
 YDGR+GE FNNR+SVGYMYMKL HHMVDKLGHARS GPYSLVTTQPLGGKAQ+GGQRPGE
 25 Sbjet: 1019 YDGRSGEAFNNRISVGYMYMKLHHMVDKLGHARS GPYSLVTTQCP+GGKAQGGQRPGE 1078

Query: 1074 MEVWALEAYGASNVLQEBILTYKSDDVTRGLKYEAITKGPVKPGVPESFVLVKEQLS 1133
 MEVWALEAYGA+ LQEBILT KSDDV GR+K YEAI KG+ +P+PGVPESF+VL+KEQLS
 Sbjet: 1079 MEVWALEAYGAYTLQEBILTYKSDDVGRVKTVEAIVKGBSVPEPGVPESFKVLVKEQLS 1138

Query: 1134 LGLDMRVLEDDNEVELRDLDEGEDDVMHVD 1166
 LG+D++L D+ R+E+RD+D DDD + +D
 30 Sbjet: 1139 LGMDEVMLSADEEETEMRDM---DDDFINQND 1168

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 379> which encodes the amino acid
 35 sequence <SEQ ID 380>. Analysis of this protein sequence reveals the following:

Possible site: 61
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 40 bacterial cytoplasm --- Certainty=0.3392 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

45 Identities = 1129/1190 (94%), Positives = 1168/1190 (97%), Gaps = 3/1190 (0%)

Query: 1 MAGHEVYQYGHKTRRSFSRIKEVLDPNLIEIQTDSQDFDLAGLKEVFDVLPISNPTD 60
 +AGHEV+YGHKTRRSFSRIKEVLDPNLIEIQTDSQDFDL+GLKEVFDVLPISNPTD
 50 Sbjet: 1 LAGHEVYQYGHKTRRSFSRIKEVLDPNLIEIQTDSQDFDLDSGLKEVFDVLPISNPTD 60

Query: 61 TMLEFVGVEKPKYKYLEEARIHDAYSAPIFVTRFLVNKETGEIKTQEVFFGDFPMT 120
 TM+LEFVGVE KPKYKYLEEARIHDAYSAPIFVTRFLVNKETGEIKTQEVFFGDFPMT
 55 Sbjet: 61 TMLEFVGVEFKPKYKYLEEARIHDAYSAPIFVTRFLVNKETGEIKTQEVFFGDFPMT 120

Query: 121 EMGTFIINGGERLIYSQVLRSPGVYPNDKVDKNGKVGYSVTIPNRGANLELTDKDLIA 180
 EMGTFIINGGERLIYSQVLRSPGVYPNDKVDKNGKVGYSVTIPNRGANLELTD+KDLIA
 55 Sbjet: 121 EMGTFIINGGERLIYSQVLRSPGVYPNDKVDKNGKVGYSVTIPNRGANLELTDKDLIA 180

Query: 181 YTRIDRTKIPFTTLVRALGFSQGDIEVDIPGDSRLVNTETKDTHNPSDSRTDEALKE 240
 YTRIDRTKIPFTTLVRALGFSQGDIEVDIPG+S+LVNRTETKDTHNPSDSRTDEALKE
 60 Sbjet: 181 YTRIDRTKIPFTTLVRALGFSQGDIEVDIPGDSRLVNTETKDTHNPSDSRTDEALKE 240

Query: 241 IYERLRPGEKPTADSSRLILARFFDPRRYDLAAGRYKINKKIMLKTRILMLTIAENIV 300
 IYERLRPGEKPTADSSRLILARFFD RYDLAAGRYK+NICKIM+KTRILMLTIAENIV
 65 Sbjet: 241 IYERLRPGEKPTADSSRLILARFFDARYDYDLAAGRYKVNCKIMLKTRILMLTIAENIV 300

5	Query: 301	DGETGEILVBAQTMTRDVIDSLAEHIDGDLNKFVYTPNDYAVVTEPVLQKFKVVAPTD 360
	Sbjct: 301	DAETGEILVBAQTMTRSVIESIEEHLDDGDLNKFVYTPNDYAVVTEPVLQKFKVVAPTD 360
10	Query: 361	PDRVVTIVGNANFDDKVRALTADILARMSYFLNLABGIGKVDIDHLEGNRIKRAVGSLL 420
	Sbjct: 361	PDRVVTIVGNANFDDKVRALTADILARMSYFLNLABGIGKVDIDHLEGNRIKRAVGSLL 420
15	Query: 421	ANQFRIGLARMEKRVREMSVQDQNVLTPOQIINIRPVTAAVGFPGSSQLSQPMQDQNP 480
	Sbjct: 421	ANQFRIGLARMEKRVREMSVQDQNVLTPOQIINIRPVTAAVGFPGSSQLSQPMQDQNP 480
20	Query: 481	LSELSHKRRLSALGPGLTRDRAGYEVDRVYTHYGRMCPLETPEGPNIGLINLSSPGH 540
	Sbjct: 481	LSELSHKRRLSALGPGLTRDRAGYEVDRVYTHYGRMCPLETPEGPNIGLINLSSPGH 540
25	Query: 541	LNKYGFICTPYRKVDRTGAVTNEIVMLTADDEEFTVAQANSKLNEDGTFABEIVMGRH 600
	Sbjct: 541	LNKYGFICTPYRKVDRTGAVTNEIVMLTADDEEFTVAQANSKLNEDGTFABEIVMGRH 600
30	Query: 601	QGNQGEFFSSIVDFVDVSFKQVAVATACIPFLENDSDRALMGANMQRAVELIDPKAP 660
	Sbjct: 601	QGNQGEFFSSIVDFVDVSFKQVAVATACIPFLENDSDRALMGANMQRAVELIDPKAP 660
35	Query: 661	YVGTMEYQAAHDSGAFLAKHDKRVI FSDAEKVEVRREDGSLDVYHVKFRFRNSGTAY 720
	Sbjct: 661	YVGTMEYQAAHDSGAFLAKHDKRVI FSDAEKVEVRREDGSLDVYHVKFRFRNSGTAY 720
40	Query: 721	NQRTLVKVGDI VEKGDFIADGPEMENGDMALQNFVVAVTWEGYNFEDAVIMSERLVKE 780
	Sbjct: 721	NQRTLVKVGDI VEKGDFIADGPEMENGDMALQNFVVAVTWEGYNFEDAVIMSERLVKE 780
45	Query: 781	DVYTSVHLEEFESSETRDTKLGPEETREI FNVGDSRLDDEMGIIRIGAEVKEGGDILVG 840
	Sbjct: 781	DVYTSVHLEEFESSETRDTKLGPEETREI FNVGDSRLDDEMGIIRIGAEVKEGGDILVG 840
50	Query: 841	KVTPGEEKDLSAERLLHATFGDKSREVRDTSLRVPHGGDGVDRVKIFTRANGDELQSG 900
	Sbjct: 841	KVTPGEEKDLSAERLLHATFGDKSREVRDTSLRVPHGGDGVDRVKIFTRANGDELQSG 900
55	Query: 901	VNMLVRVYIAKRRKIKVGDIMAGRHGNGVVSRI VPEDMFYLPGDTPVDIMNLPLGVFS 960
	Sbjct: 901	VNMLVRVYIAKRRKIKVGDIMAGRHGNGVVSRI VPEDMFYLPGDTPVDIMNLPLGVFS 960
60	Query: 961	RMNIGQVMELHGMNAARNLGIHIATPVPDGASSEDLMETVREAGMDSDAKTVLYDGRG 1020
	Sbjct: 961	RMNIGQVMELHGMNAARNLGIHIATPVPDGASSEDLMETVREAGMDSDAKTVLYDGRG 1020
65	Query: 1021	PFDNRVSVGVMYIKLHHMVDKLEHARSVGSPLVLTQQLGGQAQGGQRFGEHMEVWALE 1080
	Sbjct: 1021	PFDNRVSVGVMYIKLHHMVDKLEHARSVGSPLVLTQQLGGQAQGGQRFGEHMEVWALE 1080
70	Query: 1081	AYGASNVLQELILTYKSDVTRGLKAYRATTKGKPIPKPGVPESFRVLVKRLQSLGLDMRV 1140
	Sbjct: 1081	AYGASNVLQELILTYKSDVTRGLKAYRATTKGKPIPKPGVPESFRVLVKRLQSLGLDMRV 1140
75	Query: 1141	LQEDDNEVELEDLQEGDDDDMHVDLSEKARVKEAEKQAEQVSEVQKE 1190
	Sbjct: 1141	LQEDDNEVELEDLQEGDDDDMHVDLSEKARVKEAEKQAEQVSEVQKE 1190

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 114

A DNA sequence (GBSx0118) was identified in *S.galactiae* <SEQ ID 381> which encodes the amino acid sequence <SEQ ID 382>. This protein is predicted to be DNA-directed RNA polymerase, beta subunit (rpoC). Analysis of this protein sequence reveals the following:

```

5   Possible site: 32
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
10      bacterial cytoplasm --- Certainty=0.1892 (Affirmative) < succ>
        bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
        bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 383> which encodes the amino acid sequence <SEQ ID 384>. Analysis of this protein sequence reveals the following:

```

15   Possible site: 22
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
20      bacterial cytoplasm --- Certainty=0.2128 (Affirmative) < succ>
        bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
        bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

```

25   Identities = 1148/1205 (95%), Positives = 1177/1205 (97%)

Query: 11 VVDVNRFKSMQITLASPSKVRWSGYGEVKKPPTINRYTLKFERBGLFDEVIFGPTKWEC 70
       VVDVNRFKSMQITLASPSKVRWSGYGEVKKPPTINRYTLKFERBGLFDEVIFGPTKWEC
Sbjct: 1 VVDVNRFKSMQITLASPSKVRWSGYGEVKKPPTINRYTLKFERBGLFDEVIFGPTKWEC 60

30   Query: 71 ACGKYKRIYKGIICDRGQVEVTRAKVRERMGHIELKAPVSHIWFYKGIIPSRMGLTLM 130
       ACGKYKRIYKGIICDRGQVEVTRAKVRERMGHIELKAPVSHIWFYKGIIPSRMGLTLM
Sbjct: 61 ACGKYKRIYKGIICDRGQVEVTRAKVRERMGHIELKAPVSHIWFYKGIIPSRMGLTLM 120

35   Query: 131 SPRALEEVIYFAANYVVIDPMDTLEPKSLLTEREYREKLQYGYGSGFVAKMGAEAIQDL 190
       SPRALEEVIYFAANYVVIDPMDTLEPKSLLTEREYREKLQYGYGSGFVAKMGAEAIQDL
Sbjct: 121 SPRALEEVIYFAANYVVIDPMDTLEPKSLLTEREYREKLQYGYGSGFVAKMGAEAIQDL 180

40   Query: 191 KRVDLDAEIAVLKEELKASGQKRIKVRRLVDLDAFKSGNKPENMVNLILPVIIPDLR 250
       KRVDL AETL AKEELKSA+GOKR+KAVRRLVDLDAF KSGNKPENMVNLILPVIIPDLR
Sbjct: 181 KRVDLAAETLAEELKASGQKRIKAVRRLVDLDAFKSGNKPENMVNLILPVIIPDLR 240

45   Query: 251 FVQVLDGGRFAASDINDLYRRVIRNRNRLARLELNAPGIIVQNEKRMLEAVDALIND 310
       FVQVLDGGRFAASDINDLYRRVIRNRNRLARLELNAPGIIVQNEKRMLEAVDALIND
Sbjct: 241 FVQVLDGGRFAASDINDLYRRVIRNRNRLARLELNAPGIIVQNEKRMLEAVDALIND 300

50   Query: 311 RRGRRITGPGSRPLKSLSHMLKKGQGRFQNLIGKRVDFSGRSVIAGPTLKNYQGVPR 370
       RRGRRITGPGSRPLKSLSHMLKKGQGRFQNLIGKRVDFSGRSVIAGPTLKNYQGVPR
Sbjct: 301 RRGRRITGPGSRPLKSLSHMLKKGQGRFQNLIGKRVDFSGRSVIAGPTLKNYQGVPR 360

55   Query: 371 EMAIELFKPFVMEKIVARDLACNVKAAKGMVERGDERINDILEEVIKHPVLLNRAPTLH 430
       EMAIELFKPFVMEKIVAA++ AGNVKAAKGMVERGDERINDILEEVIKHPVLLNRAPTLH
Sbjct: 361 EMAIELFKPFVMEKIVAKEYAGNVKAAKGMVERGDERINDILEEVIKHPVLLNRAPTLH 420

60   Query: 431 RLGIQAFEPVLIDGKALRLHPIVCEAYNADPDGQMAIHVPLSEGAQAKARLLMLAAEHI 490
       RLGIQAFEPVLIDGKALRLHPIVCEAYNADPDGQMAIHVPLSEGAQAKARLLMLAAEHI
Sbjct: 421 RLGIQAFEPVLIDGKALRLHPIVCEAYNADPDGQMAIHVPLSEGAQAKARLLMLAAEHI 480

Query: 491 LNPDKGKPVVTPSQDMVLGNYYLTMDAGRGEGBMIFKDKDEAVMAYNGYVHHVTRVGI 550
Sbjct: 481 LNPDKGKPVVTPSQDMVLGNYYLTMDAGRGEGBMIFKDKDEAVMAYNRYVHHLHVRVGI 540

Query: 551 AVDSMPNKPWTERQKHKIMVTTVGKILFNDIMPDLPLYLIPFNANLTERPTDKYPLRFG 610

```

AVDSMPNKWF + Q+HKIMVTTVGKILFNDIMPEDLPYL EPNANALTS TPDKYFLEPG
 Sbjct: 541 AVDSMPNFKWQNRHKIMVTTVGKILFNDIMPEDLPYLQEPNANALTSPTDPKYFLEPG 600

Query: 611 QDIQAVINDNLEINIPFKKKNLGNIIAETFFKFRITETSFLDELKDLGYHSTLAGLTVG 670
 5 QDIQ VID L+IN+PFKKKNLGNIIAETFFKFRITETSFLDELKDLGYHSTLAGLTVG
 Sbjct: 601 QDIQSEIDRLDINVPFKKKNLGNIIAETFFKFRITETSFLDELKDLGYHSTLAGLTVG 660

Query: 671 IADIPIVDNKAEIIDAHHVRVEDINKAPRGLMTREDRYVAVITTRKRAKRALEKKRIET 730
 IADIPIVDNKAEIIDAHHVRVE+INKAPRGLMT+DRYVAVITTRKRAKRALEKKRIET
 10 Sbjct: 661 IADIPIVDNKAEIIDAHHVRVEINKAPRGLMTDDRYVAVITTRKRAKRALEKKRIET 720

Query: 731 QDPKNPIVMMDDSGARGNISNPSQLAGMRGLMAAPNGRIMELPILSNFRGLSVLEMPFS 790
 QDPKNPIVMMDDSGARGNISNPSQLAGMRGLMAAPNGRIMELPILSNFRGLSVLEMPFS
 15 Sbjct: 721 QDPKNPIVMMDDSGARGNISNPSQLAGMRGLMAAPNGRIMELPILSNFRGLSVLEMPFS 780

Query: 791 THGARKGMDTALKTADSGYLTRRLVDVAQVLIREDDCGTDRGLITAITDGKEVTSL 850
 THGARKGMDTALKTADSGYLTRRLVDVAQVLIREDDCGTDRGL IAITDGKEVTSL
 Sbjct: 781 THGARKGMDTALKTADSGYLTRRLVDVAQVLIREDDCGTDRGLLRAITDGKEVTSL 840

Query: 851 EERLIGRYTKSKHPETGEILWGADTLITEDMAKVVKAGVEEVTIRSVFTCNTRHGVC 910
 EERL GRYT+KS+KHPETGE+L+GAD LITEDMA K+V AGVEEVTIRSVFTC TRHGVC
 20 Sbjct: 841 EERLQGRYTRKSVKHPETGEVLIGADLITEDMARKIVDAGVEEVTIRSVFTCATRHGVC 900

Query: 911 RHCYGINLATGDAVEVGAEVGTIAAQSIGEPGTQLTMRFTHTGGVASNTIDTQGLPRIQE 970
 RHCYGINLATGDAVEVGAEVGTIAAQSIGEPGTQLTMRFTHTGGVASNTIDTQGLPRIQE
 25 Sbjct: 901 RHCYGINLATGDAVEVGAEVGTIAAQSIGEPGTQLTMRFTHTGGVASNTIDTQGLPRIQE 960

Query: 971 IFEARNPKGEAVITEVKGVEVAIEEDSSTRTKVFKVKGQTBGEYVVPFTARMKVEVGDE 1030
 IFEARNPKGEAVITEVKG VV IEED+STRTKV+V+G+TG GEYV+PFTARMKVEVGDE
 30 Sbjct: 961 IFEARNPKGEAVITEVKGNNVVEIEEDASTRTKVVQGTGMGEYVLPFTARMKVEVGDE 1020

Query: 1031 VARGAALTGSGIQPKRLLEVRDTLSVETYLLEAVQKVYRSGQVEIGDKHVEVMVRQMLRK 1090
 V RGAALTGSGIQPKRLLEVRDTLSVETYLLEAVQKVYRSGQVEIGDKHVEVMVRQMLRK
 35 Sbjct: 1021 VNRGAALTGSGIQPKRLLEVRDTLSVETYLLEAVQKVYRSGQVEIGDKHVEVMVRQMLRK 1080

Query: 1091 VRVMDPGDTDLLPGLTMDISDFTDANKDIVISGGIPATSRPVLMSGITKASLETNSPLSAA 1150
 VRVMDPGDTDLLPGLTMDISDFTDANKDIVISGGIPATSRPVLMSGITKASLETNSPLSAA
 Sbjct: 1081 VRVMDPGDTDLLPGLTMDISDFTDANKDIVISGGIPATSRPVLMSGITKASLETNSPLSAA 1140

Query: 1151 SFQETTRVLTDAAIRGKKDHLGLKENVIIGKIIPAGTGARYRNIEPLAVNEVEIIGT 1210
 SFQETTRVLTDAAIRGKKDHLGLKENVIIGKIIPAGTGARYRNIEP A+NE+E+I+ T
 40 Sbjct: 1141 SFQETTRVLTDAAIRGKKDHLGLKENVIIGKIIPAGTGARYRNIEPQWNEVEIIGT 1200

Query: 1211 PVDAE 1215
 V AE
 45 Sbjct: 1201 EVSAE 1205

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

50 Example 115

A DNA sequence (GBSx0120) was identified in *S. agalactiae* <SEQ ID 385> which encodes the amino acid sequence <SEQ ID 386>. This protein is predicted to be a DNA binding protein. Analysis of this protein sequence reveals the following:

Possible site: 19
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.4727(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 60 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

-188-

>GP:AAC45309 GB:U81957 putative DNA binding protein [Streptococcus gordonii]
Identities = 42/99 (42%), Positives = 75/99 (75%)

5 Query: 1 MYQVVKMGDWEPMWFI B GWEEDITEIAEYDITLSEALLYPQERMDRGQEKWPYQSKSSI 60
MY+VV+M+GD+EPWPF+GWE DI + + +AL+++W+ + + + +S+L
Sbjct: 1 MYRVVMYGDPEWPFIDGWEENDIQORPEKYDALKFYIKQMLKLETPEKYEKRSDI 60

Query: 61 LATFWSIKRRKRCDEYLQQYHSLMLKEWQIPKEE 99
+ FW+ ++RWCECD+Y+QY S+LL++ + IPK +
10 Sbjct: 61 MIVFWNENDQKRCEDDYVQYRSIILLKKEKVIKPSK 99

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 387> which encodes the amino acid sequence <SEQ ID 388>. Analysis of this protein sequence reveals the following:

15 Possible site: 36
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.4741 (Affirmative) < succ>
20 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 61/121 (50%), Positives = 83/121 (68%)

25 Query: 1 MYQVVKMGDWEPMWFI R GWEEDITEIAEYDITLSEALLYPQERMDRGQEKWPYQSKSSI 60
MYQV+IM+GDWEPMWFI+G++DI + + +EAL YF +EW R + +P+ S+ +L
Sbjct: 1 MYQVIMKGDWEPMWFI D GQDDIIDQPSDWQFALDYFNCWQRKKAIFPSYHSKML 60

30 Query: 61 LATFWSIKRRKRCDEYLQQYHSLMLKRWQIPKEESIERPFVFNKIAKLPASACIANL 121
LATFW ++RWCE+CDE LQQ+HSL+LLK +P I FE N ++ C IANL
Sbjct: 61 LATFWKEKDRKRCDEEDLQPFHSLLLKNDIVESNNTYIPEFQRNDSQFVYALCKIANL 121

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

35 Example 116

A DNA sequence (GBSx0121) was identified in *S.galactiae* <SEQ ID 389> which encodes the amino acid sequence <SEQ ID 390>. Analysis of this protein sequence reveals the following:

40 Possible site: 18
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.2433 (Affirmative) < succ>
45 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAC45310 GB:U81957 putative ABC transporter subunit ComYA
[Streptococcus gordonii]
Identities = 203/319 (63%), Positives = 255/319 (79%), Gaps = 1/319 (0%)

50 Query: 1 MVQSLAQVHQVAVFVNAQDIYLIIPKGD CYELNMRIDERRFDIVFPFNKASLI SHKFF 60
MVQ +A+ ++ QA E AQDIY +PK DCEYNMREI DERRFI ++P++A++ISHKFF
Sbjct: 1 MVQKLAQVIRQAQRECAQDIYFVPKND CYELNMRIGERRFIQTYDFQLAVALISHKFF 60

55 Query: 61 VAGMNVGEKRRSQSGCDYRLSEGRISVLSRLSSVGDYRQGSILVIRLYSHGQDLKWF 120
+AGMNVGEKRRSQSGCDY ++ S+RLS+VGDYRG ESLVIR+L+ +L+WF
Sbjct: 61 LAGMNVGEKRRSQSGCDYRYDD-KETSI RLS+VGDYRG ESLVIR+LHDESETEIKWF 119

Query: 121 NIKQKMEVLGIRGLYLFSGPVGSGKTYLMYQLASEVFNKQIITIEDPVKIKNDKMLQIQ 180

+ +++E RGLYLSPGPGSGKTTLM+QLA FK +Q+++IEDPVEIK + MLQLQ
 Sbjet: 120 HFFPRIRKKFKDRGLYLSPGPGSGSKTTLMHQLAQLKPKGQCVMSIEDPVEIKQRMQLQLQ 179
 Query: 181 LNEIDIGMTYDALIKLSLRHRPDILIGEIRDQATARAVIRASLTGMVFSITHAKSIPGV 240
 5 LNE IG+TY++LILKLSLRHRPD+LIGEIRD TARAV+RASLTG VFSTHAKSIPGV
 Sbjet: 180 LNETIGLTYESILIKLSLRHRPDILLIGEIRDSRTARAVRASLTGATVFSTHAKSIPGV 239
 Query: 241 YDRLELGVNYQELNSLKLIAVQRLIGGGSILDFETGNPKHSSDKWNRQVDILAREGH 300
 Y+RL+ELGV+ +EL+ L+ I YQRLIGG +IDF + N+++H WN+Q+D L GH
 10 Sbjet: 240 YERLLELGVSEELKIVLGICVQRLIGGGVIDFASDNYQEHPTVWQQIDQLLAAGH 299
 Query: 301 ISKKQAQVEKIIPOETTES 319
 I +QA+ EKI Q+ S
 15 Sbjet: 300 IHPQAEAEKIRNQAKTS 318

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 391> which encodes the amino acid sequence <SEQ ID 392>. Analysis of this protein sequence reveals the following:

Possible site: 18
 >>> Seems to have no N-terminal signal sequence
 20 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1846 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 25 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 207/312 (66%), Positives = 257/312 (82%)
 Query: 1 MVQSLAKQVHQAVEVNAQDIYIIPKQDCYELVMRIDDERFIDVFEFNRASLISHFKF 60
 30 MVQ+LAK ++ +A +V+AQDIYI+P+ D Y+L++RI DERR +DV++ +RMA LISHFKF
 Sbjet: 1 MVQALAKAILAKABQVHAQDIYIIPRACQYDLFLRIGDERRLDVYVQSDRMAPLSIFKFK 60
 Query: 61 VAGMIVGEKRRSQGLSCDYELSSGRVLSRLSSVGDYRGQBSLIRILYSGHQDLKWFND 120
 VAGM VGEKRR Q+GSCDY+LS+ + +SLRLSSVGDYRGQBSLIVIR+L+ ++ + YWFD
 35 Sbjet: 61 VAGMIVGEKRRQVGSQCDYKLSNDKQLSLRLSSVGDYRGQBSLIRILHQNKSVHYWFD 120
 Query: 121 NIKQMKVEVLQIRGLYLPSPGPGSGKTTLMYQLASEVFNKQIITIEDPVEIKNDRLQLQ 180
 + ++ +G RGLYL+FPVPGSGKTTLMYQL S + Q+I+IEDPVEIKN ++LQLQ
 40 Sbjet: 121 GLTKVANQVGGRLYLFPAGVPGSGKTTLMYQLISNYHQEAVISIEDPVEIKNHQILQLQ 180
 Query: 181 LNEIDIGMTYDALIKLSLRHRPDILIGEIRDQATARAVIRASLTGMVFSITHAKSIPGV 240
 +N+DIGMTYD LILKLSLRHRPDIL+IGEIRD TARAVIRASLTG MVFST+HAKSI GV
 45 Sbjet: 181 VNDDIGMTYDNLILKLSLRHRPDILLIGEIRDSRTARAVIRASLTGMVFSITHAKSISGV 240
 Query: 241 YDRLELGVNYQELNSLKLIAVQRLIGGGSILDFETGNPKHSSDKWNRQVDILAREGH 300
 Y RL+ELGV EL N L LIAVQRL+ GG-LID F+ +SS WN+Q+D L E GH
 50 Sbjet: 241 YARLLELGVTKAZLSNCLALIAVQRLINGGALIDSTQNEFETYSNNWQQIDQLLEAGH 300
 Query: 301 ISKKQAQVEKII 312
 ++ QA++EKII
 Sbjet: 301 LNPRQAKLEKII 312

SEQ ID 390 (GBS63) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 5 (lane 5; MW 39kDa). It was also expressed in *E.coli* as a GST-fusion product.

55 SDS-PAGE analysis of total cell extract is shown in Figure 13 (lane 2; MW 64kDa).

The GBS63-GST fusion product was purified (Figure 101A; see also Figure 191, lane 3) and used to immunise mice (lane 1 product; 20µg/mouse). The resulting antiserum was used for Western blot (Figure 101B), FACS (Figure 101C), and in the *in vivo* passive protection assay (Table III). These tests confirm that the protein is immunoaccessible on GBS bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 117

A DNA sequence (GBSx0122) was identified in *S.galactiae* <SEQ ID 393> which encodes the amino acid sequence <SEQ ID 394>. This protein is predicted to be competence protein (mshG). Analysis of this protein sequence reveals the following:

```

Possible site: 49
>>> Seems to have no N-terminal signal sequence
10  INTEGRAL    Likelihood =-14.65    Transmembrane    123 - 139 ( 113 - 144)
    INTEGRAL    Likelihood =-13.53    Transmembrane    272 - 288 ( 264 - 295)
    INTEGRAL    Likelihood = -8.55    Transmembrane    79 - 95 ( 75 - 102)
    INTEGRAL    Likelihood = -0.00    Transmembrane    146 - 162 ( 146 - 162)

----- Final Results -----
15  bacterial membrane --- Certainty=0.6859(Affirmative) < succ>
    bacterial outside --- Certainty=0.0000(Not Clear) < succ>
    bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 9489> which encodes amino acid sequence <SEQ ID 9490> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AAC45311 GB:U81957 putative ABC transporter subunit ComYB
[Streptococcus gordonii]
Identities = 161/280 (57%), Positives = 219/280 (77%)

25  Query: 19  MNKALLECKDLKMLGELGFSDTVITQVALADLHGNISRLKTKTESVLANLLVRKKVIE 78
    M + L G+ S+++ LGFSD V+TQ++LA+LHGN+S +LLKTE YL NL V+KK+IE
    Sbjct: 1  MRQGLANGQAFSEIMASLGFSDAVVTQLSLAELHGNLSALLKIREVLDNLAKVKKKLE 60

30  Query: 79  VATYPLILLSPFLVLMIGLERNYLMPOLGENNFAETLITNVFNIFLLLLAVLIFSLIFYI 138
    VATYP++LL FLVLMIGLERNYL+POL NFAT+LI ++F IFLL + ++L + Y+
    Sbjct: 61  VATYPMMLLGFLVLMIGLERNYLLPOLSSQNFAATQLIGHFTIFLLTVMLGLLIGATYL 120

35  Query: 139  IQKRLSRIKVACFLTTIPNGSYVKLYLTATYAREWGNLLSGGLELDQIVKVMONQSKL 198
    + K RI V FL + P VGS+V++YLTATYAREWGN++ QG+EL QI ++MQ Q+S L
    Sbjct: 121  VFKQKQKIPVYSFLARLPFVGSFVRIYLTATYAREWGNNGQGLELSQIFQIMQBQBSVL 180

40  Query: 199  FREITGYDMEEGFLSGKAFHQKVLDPYFFLTELSELMIEYGVKAKIGTELDTYADEKVEDF 258
    F+RIG D+ + +G+ F K+ YPPF ELSL+IEYG+VE+KLG+EL+IYA + WE+F
    Sbjct: 181  FQRTGQDLQALNQGRFSDKIASYPFFKKLSLITREYGEVSKLSGLSELTALKTWEEF 240

    Query: 259  FTKIARATQLIQPVIFIPVALIIVMIYAAMLLPYQNMMEI 298
    F + R LIQP++F+VAL+IV++YAAMLLP+YQNMME+
    Sbjct: 241  FGRVNRITMNLIQPLVVFVVALMIVLLYAAMLLPYQNMMEV 280

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 395> which encodes the amino acid sequence <SEQ ID 396>. Analysis of this protein sequence reveals the following:

```

Possible site: 43
>>> Seems to have no N-terminal signal sequence
50  INTEGRAL    Likelihood =-12.52    Transmembrane    317 - 333 ( 309 - 339)
    INTEGRAL    Likelihood =-10.14    Transmembrane    123 - 139 ( 119 - 147)
    INTEGRAL    Likelihood = -6.95    Transmembrane    164 - 180 ( 161 - 183)

----- Final Results -----
55  bacterial membrane --- Certainty=0.6010(Affirmative) < succ>
    bacterial outside --- Certainty=0.0000(Not Clear) < succ>
    bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the databases:

>GP:ARC45311 GB:U81957 putative ABC transporter subunit ComYB
[Streptococcus gordonii]
Identities = 139/278 (50%), Positives = 207/278 (74%)

Query: 63 MEESLLKQGLAMLSGLGSPDAILTQISLADRHGNIETTLVAIQHYLNQMARIRKKTVE 122
M + L GQ +++++ LGFSDA++TQ+SLA+ EGN+ L+ I+ YL+ +A++K +E
Sbjct: 1 MRQLANQCAFSEIMAGLGFSDAVVTQLSLAEHLGNLSALLKITEYLDNLAKVKKLIE 60

Query: 123 VITYPLILLLLFLVMMGLRRYLVPQLETNQNTTYFLNHPFAFPGCGILLLPGMVL 182
V TYP++LL FL ++M+GLR YL+PQL +QN T + H P F+ L+ L G ++L
Sbjct: 61 VATYPMMLLGLFVLMIGLRNLYLLPQLSQCNPATQLIGHLPTIFLLVLMMLGLNGNLY 120

Query: 183 RWRQSRLKLYSRLSRYPFLGKLLKQVLTYSYAREWGTLIGQGLDMLTILDIMAEKSSL 242
++ Q R+ +YS L+R PF+G ++ YLT+YYAREWG +IGQL+L I IM ++S L
Sbjct: 121 VFKQKRIPVYSFLARLPFVGSPVRIYLTAYAREWGNMIGQLELSQIFQIMQBRQSVL 180

Query: 243 MGELEADIRMSLLESGQAFHIKQVATYPPFKKELSLMIEYGEIKSGKLEIYAQBSWEQF 302
+E+ +D+ +L GQ P K+A+YPPFKKELSL+IEYGE+KSLG+ELEIYA ++WE+F
Sbjct: 181 FQEGDQLGQALNQGEFSKDIASYPFFKGLSLIIEYGEVKSGLSELEIYALKTWEEF 240

Query: 303 PSQLYQVTLQIQPAIFLVVAVTIWMIYAAILLPIYQNM 340
F ++ + LIQP +F+ VA+ IV++YAA+LLP+YQNM
Sbjct: 241 FGRVNETNLIQPLVFPVFMVLMVLLYAMLLPIYQNM 278

An alignment of the GAS and GBS proteins is shown below:

Identities = 148/297 (49%), Positives = 209/297 (69%), Gaps = 2/297 (0%)

Query: 1 MVTFLKRSKLLSDCYTDSMNKALLEGKDLKXMGELGSPDVTITQVALADLHGNISRL 60
++ FLKRS+LL Y M ++LL+G+ L+ ML LGFSD ++TQ++LAD HGNI +L+
Sbjct: 45 VIAFLKRSQQLQDLVYLMMEESLLKQGLAMLSGLGSPDAILTQISLADRHGNIETTLV 104

Query: 61 KIESYLANLLLVKRVKIEVATYPLILLSFLVLMIGLRNTLMPQLENNFATRLTITVFN 120
I+ YL + +R+K +EV TYPLILL FL ++M+GLR YL+PQL N T + + P
Sbjct: 105 AIQHYLNQMARIRKKTVEVITYPLILLLLFLVMMGLRRYLVPQLETNQNTTYFLNHPFA 164

Query: 121 IFL-LLAVVLIFSLIFYIIQKRLSRKIVACFLITIPLVGSYVGLYLTAYAREWGILL 179
F+ ++L+F ++ ++ + SR+K+ L+ P +G +K YLT+YYAREWG L+
Sbjct: 165 FFIGFCSGLILLFGMV-WLRWRQSRLKLYSRLSRYPFLGKLLKQVLTYSYAREWGTLIG 223

Query: 180 QGIELDQIVKVMNQKSLFREIGYDMEBGFSGKAFHQKVLDPYFFLITSLMIEYGV 239
QG++L I+ +M +KS L+E+ D+ L G+APH KV YPFF ELSLMIEYG+
Sbjct: 224 QGLDMLTILDIMAEKSSIMKELADIRMSLLESGQAFHIKQVATYPPFKKELSLMIEYGEI 283

Query: 240 KAKLTGTELDIYADEKWEDEFKTLARATLIQVPIFVFIYVALIIVMIYAAILLPIYQNM 296
K+KLG EL+IYA E WE FF++L + TQLIQ IF+ VA+ IVMYIAA+LLP+YQNM
Sbjct: 284 KSKLGAELEIYACBSWEQFPSQLYQVTLQIQPAIFLVVAVTIWMIYAAILLPIYQNM 340

A related GBS gene <SEQ ID 8493> and protein <SEQ ID 8494> were also identified. Analysis of this protein sequence reveals the following:

Lipop: Possible site: -1 Crend: 9
SRCFLG: 0
McG: Length of UR: 2
Peak Value of UR: 1.24
Net Charge of CR: 0
McG: Discrim Score: -8.94
GVH: Signal Score (-7.5): -4.08
Possible site: 31
>>> Seems to have no N-terminal signal sequence
Amino Acid Composition: calculated from 1
ALOM program count: 4 value: -14.65 threshold: 0.0

INTEGRAL	Likelihood = -14.65	Transmembrane	105 - 121 (95 - 126)
INTEGRAL	Likelihood = -13.53	Transmembrane	254 - 270 (246 - 277)
INTEGRAL	Likelihood = -8.55	Transmembrane	61 - 77 (57 - 84)

```
PERIPHERAL Likelihood = 5.09      14
modified ALOM score: 3.43
icml HYPRID: 7 CFP: 0.686
```

5 *** Reasoning Step: 3

----- Final Results -----

```
bacterial membrane --- Certainty=0.6859(Affirmative) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>
```

The protein has homology with the following sequences in the databases:

57.5/79.7% over 279aa

15 GP|2058545| putative ABC transporter subunit ComYB Insert characterized Streptococcus gordonii

ORF00008 (355 - 1194 of 1500)

GP|2058545|gb|AAC45311.1||U01957(1 - 280 of 282) putative ABC transporter subunit ComYB
{Streptococcus gordonii}

20 {Streptococcus
%Match = 33.8

\$Identity = 57.5 \$Similarity = 79.6

Matches = 161 Mismatches = 57 Conservative Sub.s = 62

25 144 174 204 234 264 294 324 354
TLROVILKYNTHOTSGIDKWISSLKKDISVRNRHKSLLKLRKQKVVOLFNNLFASGFSLTDMVTFLKRSKLLSDCYTDS

384 414 444 474 504 534 564 594
MNKALLEGKDLKSMELGELPSTDVITQVALADLHGNISRLSKIESYLANILLVRKKIVLEVATYPLILSLFVLIMIGLR
| | | | | : : : : : | : : : : : | : : : : : | : : : : : | : : : : : | : : : : : | : : : : : | : : : : :
30 MRQGLANGAQA FSEIMASGELSGDAVVQSLSLALHGNLSLALKITEEYLDNLAKKKLIVLEVATYPMWLLFVLIMIGLR
10 20 30 40 50 60 70 80

[illegible]

40 864 894 924 954 984 1014 1044 1074
SQGIELDIQVVMQNQSKLFRREIGYDMEBGLFSGKAFHQVLVDYPFFITLSLMIEYGQVKAKGLGTEDLYADEKWEDF
||| || | : :: || | : || | : : : || | : || | ||||| : ||| : ||| : || : || : ||
GGQLSELISQIQINQRORSVLFQOEGIDGLQALQNGSFSDIASPYPFKKELSLIIRYGEVKSCLKSELEILYALKTWEFF
 170 200 230 260

[illegible]

50

SEQ ID 8494 (GBS49) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 11 (lane 5; MW 15kDa). It was also expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 15 (lane 5; MW 60kDa).

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 118

A DNA sequence (GBSx0123) was identified in *S.agalactiae* <SEQ ID 397> which encodes the amino acid sequence <SEQ ID 398>. This protein is predicted to be ComYD or ComGD. Analysis of this protein sequence reveals the following:

Possible site: 55
 >>> Seems to have a cleavable N-term signal seq.

----- Final Results -----

- 5 bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

- 10 >GP:CAA75315 GB:Y15043 homology to ComYD from *Streptococcus gordonii*,
 and ComGD from *Bacillus subtilis* [*Lactococcus lactis* subsp. *cremoris*]
 Identities = 56/138 (40%), Positives = 92/138 (66%), Gaps = 2/138 (1%)
- 15 Query: 12 KVKAPTLLLECLVALVITITGALLVYQGLTKLLAQQIVVMSSSQSEWVLLTQQLNARFEGAR 71
 K+APTLLLECLVAL+ I+G+LV GLT++ +Q+ + +S+ +W+ +Q+ +E GA
 Sbjct: 13 KIRAPTLLLECLVALLAISGSLVVISGLTRMIEQMKISQNDNRKDWQIFCRQMRSELGSA 72
- Query: 72 HLEYLRQNKLYLRKQKIVTFPGKSNKDDFRKTYDGRGYQPMVYGLNQCMSCTKSMVKL 131
 L+ +QN LV+ K DK + PG DDFRK+ G+GYQPM+Y L ++ +++K+
 20 Sbjct: 73 KLDNVNQNLVYTK-DKKLRFLVG-DDFRKSDRGQGYQPMVLDLKGAKIQARENLIK 130
- Query: 132 VFYFKDGLKRTFFYPKE 149
 F +G +R F Y F +
 25 Sbjct: 131 TIDFNGGERVFIYRFTD 148

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 399> which encodes the amino acid sequence <SEQ ID 400>. Analysis of this protein sequence reveals the following:

Possible site: 26

- 30 >>> Seems to have a cleavable N-term signal seq.
- Final Results -----
- 35 bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

- 40 >GP:CAA75315 GB:Y15043 homology to ComYD from *Streptococcus gordonii*,
 and ComGD from *Bacillus subtilis* [*Lactococcus lactis* subsp. *cremoris*]
 Identities = 65/137 (47%), Positives = 84/137 (60%), Gaps = 2/137 (1%)
- Query: 8 IKAPTLLLEALIALVISGSLVYQGLTRTLKHSYLARHDQDNWLLFSHQLRRELSGAR 67
 I+APTLLLE L+ALL IGG+LV GLTR + + +W +F Q+R ELGGA+
 45 Sbjct: 14 IRAPTLLLECLVALLAISGSLVVISGLTRMIEQMKISQNDNRKDWQIFCRQMRSELGSAK 73
- Query: 68 FYKVDNKLIVVEGKKVLAPGQFKSHDFRKSAASNGKYQPMVFGISRSHTIHQSQICIT 127
 V N LVV K KK L PG DFRKS G+GYQPM+ + + I E+ I IT
 50 Sbjct: 74 LDNVNQNLVYTKDKK-LRFG-LVGDGDFRKSDRGQGYQPMVLDLKGAKIQARENLIK 131
- Query: 128 LKWKSGLERTFYAFQD 144
 + + +G ER F Y F D
 55 Sbjct: 132 TIDFNGGERVFIYRFTD 148

An alignment of the GAS and GBS proteins is shown below:

- 55 Identities = 58/137 (42%), Positives = 88/137 (63%)
- Query: 13 VKAPTLLLECLVALVITITGALLVYQGLTKLLAQQIVVMSSSQSEWVLLTQQLNARFEGAR 72
 +KAPTLLLE L+AL+ I+G+LVYQGLT+ L + ++ Q W+L +QL E GA
 60 Sbjct: 8 IKAPTLLLEALIALVISGSLVYQGLTRTLKHSYLARHDQDNWLLFSHQLRRELSGAR 67
- Query: 73 LEYLRQNKLYLRKQKIVTFPGKSNKDDFRKTYDGRGYQPMVYGLNQCMSCTKSMVKL 132
 + NKLY+ K K+ + PG+ DFRK+ +G+GYQPM+G+ + +S + +
 Sbjct: 68 FYKVDNKLIVVEGKKVLAPGQFKSHDFRKSAASNGKYQPMVFGISRSHTIHQSQICIT 127

Query: 133 FYFKDIGLKRTFYFDK 149
+K GL+RTFYY F++
Sbjct: 128 LKWKSGLERTFYFAFD 144

A related GBS gene <SEQ ID 8495> and protein <SEQ ID 8496> were also identified. Analysis of this protein sequence reveals the following:

```

Llipop: Possible site: -1      Crend: 10
McG: Discrim Score:          4.86
GVH: Signal Score (-7.5): -0.22
    Possible site: 55
>>> Seems to have a cleavable N-term signal seq.
ALON program count: 0 value: 12.47 threshold: 0.0
PERIPHERAL Likelihood = 12.47    127
modified ALON score: -2.99

*** Reasoning Step: 3

```

```

----- Final Results -----
      bacterial outside --- Certainty=0.3000(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the databases:

GP|3287181| homology to ComYD from *Streptococcus gordonii*, and ComGD from *Bacillus subtilis* {*Lactococcus lactis* subsp. *cremoris*} Inse
rt characterized

```

GRF00009(334 - 747 of 1053)
GP[3287181]emb[CAA75315.1][Y15043(13 - 148 of 150) homology to ComYD from Streptococcus
Gordonii, and ComGD from Bacillus subtilis [n
actococcus lactis subsp. cremoris]
%Match = 15.9
%Identity = 40.6 %Similarity = 68.1
Matches = 56 Mismatches = 42 Conservative Sub.s = 38

```

[illegible]

SEQ ID 398 (GBS6) was expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 1 (lane 2; MW 40kDa). It was also expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 2 (lane 2; MW 15kDa). The GBS6-GST fusion product was purified (Figure 189, lane 2) and used to immunise mice. The resulting antiserum was used for FACS (Figure 260), which confirmed that the protein is immunoreactive on GBS bacteria.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 119

A DNA sequence (GBSx0124) was identified in *S.agalactiae* <SEQ ID 401> which encodes the amino acid sequence <SEQ ID 402>. Analysis of this protein sequence reveals the following:

```
Possible site: 43
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3831 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

The protein has homology with the following sequences in the GENPEPT database:

```
>GP: AAC00317 GB: AF008220 YtxK [Bacillus subtilis]
Identities = 106/329 (32%), Positives = 176/329 (53%), Gaps = 17/329 (5%)

Query: 1 MNFEKIEAYELILENIQTIERNLKTHIYDALIEQNSYLLGSSCDLMDVVNNQKLRQLD 60
M + + YEL+ E I+N+L+ +AL E Y D + + +QK +QL
Sbjct: 1 MQKHGVHAYVELLENAIMINIGELQISYIALAEAGEMYFLKTD-QLKLPAQDKTKQLQ 59

Query: 61 LSQE-----EW-RRTFQFIFIKSAQTEQLQANHQTPTDSIGFILLFLLEE-LTSQE 109
E EW R+ FQ +K + + N Q TPD+IG + +L+ + + ++
Sbjct: 60 ALLEKAEFGTYEHWKAFQLAVLQGMK-DISHPNRQNTPTDTGLFISYLVNKPMDKK 118

Query: 110 TVDVLIEIGSGTENLAQTLLNN-SSEKLVNMGIEVDLIDLDSASTARIIGSSAQFQEDA 168
+ +L+ GTGNL T+LN S K N GIE+DD+L+ ++ + A + + + +D+
Sbjct: 119 ELTILDPALETGNLLFTVLNQLSEKTANSFGIEIDVLLKIAVAQANLLKKELELPHQDS 178

Query: 169 VRPQILKESDVIIISDLFVGYPNDGIKRYAVSSSKEHTYAHHLMEQSLKYKKKGDAI 228
+ P + D + I DLFVGYPND A+ + + + + H++AHLL +EQS+K+ K G
Sbjct: 179 LEFLFIDPVDITICDLFVGYPNDGEGARAFELKADGHSFAHLLFTEQSVRHTKFGGYLF 238

Query: 229 FLAPENLITSPQSDLLKENLKGADVIATVLEETIFGSEQNAKSI FVLKKQAEQKF--- 285
F+ P +L S QS LK++ K + A+L LP++IF +AKSI VL+RQ E
Sbjct: 239 FMIPNHLFESSQSGKLKQFFKDKVHINALIQLPKSIFKDEBARAKSILVLRQKGTGKAPG 298

Query: 286 ETFVYPLTDLQNRNMNFIENPQKWSR 314
+ + L N++ M + + F +N ++
Sbjct: 299 QILLANLPSFSNQIQLMDMAQFDEWFKK 327
```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 403> which encodes the amino acid sequence <SEQ ID 404>. Analysis of this protein sequence reveals the following:

```
Possible site: 57
>>> Seems to have an uncleavable N-term signal seq

----- Final Results -----
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
```

An alignment of the GAS and GBS proteins is shown below:

```
Identities = 223/315 (70%), Positives = 270/315 (84%)

Query: 1 MNFEKIEAYELILENIQTIERNLKTHIYDALIEQNSYLLGSSCDLMDVVNNQKLRQLD 60
M FSEKIE AY+L+LEN Q IEN LKTHIYDA++BONS+YLG+ V N+ KL+ L
Sbjct: 16 MTFEKEIEAYQLLENCQLLENDLKTHIYDAIVBONSFYLGAEASFPQVQNSDLKALC 75

Query: 61 LSQEERWRTQFIFIKSAQTEQLQANHQTPTDSIGFILLFLLEE/TSQETVDVLEIGSGT 120
```


L++ESWR+ +QF+FIK+AQTEQLQANHQFTPD+IGFILL+LLE+L+ +++++VLEIGSGT
 Sbjct: 76 LTKESWRKAYQFLFKNAQTEQLQANHQFTPD+IGFILL+LLEQLSDKSDSLEVLEIGSGT 135
 Query: 121 GNLACTLLNNSSKELNMGIEVDDLLIDLASIAETIGSSAQFIQEDAVRPQILKESDVI 180
 5 GNLACTLLNN+SK L+Y+GIE+DDLLIDLASIAE+ SSA FIQEDAVRPQ+LKESD++
 Sbjct: 136 GNLACTLLNNYSKLDYVGIELODILLIDLASIAETIMDSSAHFIQEDAVRPQLLKESDVI 195
 Query: 181 ISDLFVVOYYPNDQIAKRYAVSSSKEHTYAHHLMBQSLKYLKKDGAIFLAPENLITSPQ 240
 ISDLFVVOYYPND IAKRY V+SS +HTYAHHLMBQSLKYLKKDGAIFLAP NLITSPQ
 10 Sbjct: 196 ISDLFVVOYYPNDQIAKRYKVASSDKHTYAHHLMBQSLKYLKKDGAIFLAPVNLITSPQ 255
 Query: 241 SDLLKSWLKGYDAVIAVLTPETIPGSRQNAKSIFVLKKQAEQKPTFPVPIPLQNRN 300
 S ILK+WLK YA V+ ++TLP++IFG NAKSI VL+KQ + ETPVVP+ DL+ EN
 15 Sbjct: 256 SQLLKQWLKDYAVVITLTPDSIFGHPNNAKSIIVLQKQTHMPTFPVPIPLQKLARN 315
 Query: 301 MANFIENPQKWSRN 315
 + +F+ENF+KW N
 Sbjct: 316 IHDPMENFKWKLEN 330
 20 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 120

A DNA sequence (GBSx0125) was identified in *S. agalactiae* <SEQ ID 405> which encodes the amino acid sequence <SEQ ID 406>. This protein is predicted to be acetate kinase (ackA-1). Analysis of this protein
 25 sequence reveals the following:

Possible site: 15
 >> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 30 bacterial cytoplasm --- Certainty=0.2384 (Affixmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

35 >GP: AAC36857 GB: L17320 acetate kinase [Bacillus subtilis]
 Identities = 223/395 (56%), Positives = 293/395 (73%), gaps = 3/395 (0%)
 Query: 1 MSKTIADINAGSSSLGQQLYBHPREKVVAKGIIERIGLKDISTVKFDKKDBQILDIVDH 60
 40 MSK IADINAGSSSLK+QL+BMP E V+ KG++ERIG+ DS+ T+ + +K+ ++ DI DH
 Sbjct: 1 MSKTIADINAGSSSLKFLBHPSETVLTGKLVVERIGIADSVPTISVNGEHTVTDIDPH 60
 Query: 61 TQAVKILLEDLTAKIGI IKD PNEITGVGHVVGGEYFKESALVDDKVVQVBEELSALAPL 120
 AVK+LL LT+ GIID NRI G+GHRVV GGE F+S L+ D+ +++E++S LAPL
 45 Sbjct: 61 AVAKMLNLKLTPEGLIKDLNEIDGIGHRVVHGGKFPDSVLLTDETIKEDISELAPL 120
 Query: 121 HNPAAGIRAPREILPDITSVCVFDTAFHTMQPHLYLPPIKQYTDYKVRKYAGHT 180
 HNPA GI+AF+E+L++ +V VDTAFH TM +YLY +P +YI + +RKYG HGT
 50 Sbjct: 121 HNPAIVGIAKAFKVLPMVAVAVDTAFHQTMPQSYLYSLFVEYYEYKGIKRYGPHGT 180
 Query: 181 SHQYVAQENAKQLRPLEELKLITAHVGVGVSITANYHQQSIDTSMGFTPLAGPMGTRS 240
 SH+YV + AA+ LGREL++L+LI+ H+GNG SI A G+SIDTSMGFTPLAG MGTRS
 55 Sbjct: 181 SHKYVTERAAEILGRFLKDLRLISCHLNGASINAVBQGSIDTSMGFTPLAGVAMGTRS 240
 Query: 241 GNDIPAIIPYLVNDPELEDAAVVNMINKQSGLLGVSGTSSDMRDIAGLQSKDFNAVL 300
 G+IDPA+IPY++ + D V+N LNK+SGLAG+SG SSD+RDI + + A
 55 Sbjct: 241 GNDIPALIPYIMKTSQTD--EVLNITLNKKSGLGIGSPSSIDRDIYEATKGBEAST 298
 Query: 301 AYNVPIDRIKKFIGQYLAIVNGADAIIPTAGNGENAPLMRQDVIAGLSWPGIADPE-KN 359
 A VP RI K+IG Y A ++G DAIIIFTAG+QEN+ +R+ V+ GL+ G+ DP N
 60 Sbjct: 299 ALZVFASRIHKYIGSYAARMSGVDAIIPTAGIGNSVEVRRERVLGGLRNGVYWDPALNN 358

Query: 360 VFGYGDITKPKSKVKVLVIPTDEELMIARDVRL 394
 V G I+ P S VKV+I PTDE+MIARDV RL
 Sbjct: 359 VRGEAFISYPHSPVKVMIIPTDEEVMIAADVRL 393

- 5 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 407> which encodes the amino acid sequence <SEQ ID 408>. Analysis of this protein sequence reveals the following:

Possible site: 28

>>> Seems to have no N-terminal signal sequence
 10 INTEGRAL Likelihood = -0.22 Transmembrane 63 - 79 (63 - 79)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.1086(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 15 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP: AAC36857 GB:L17320 acetate kinase [Bacillus subtilis]
 Identities = 218/395 (55%), Positives = 293/395 (73%), Gaps = 3/395 (0%)
 20 Query: 1 MSKTIAINAGSSSLKWLQYMPPEAVLAQGIIRIGLAKDSISTVKYDGKKSEQLIDIH 60
 MSK IAINAGSSSLK+QL+MP E VL +G+ERIG+ DS+ T+ +G+K ++ DI DH
 Sbjct: 1 MSKIIAINAGSSSLKQLQFEMPEETVLTKGLVERIGIADSVFTISVGNKTEVTDIPDH 60
 25 Query: 61 TEAVKILLNDLHFGIIAAYDEITGVGHRVAGGELFKESVVDKVLQIEELSLVLAFL 120
 AVK+LLN L PGII +EI G+GHRVV GSE F +SV++ D+ +++IE++S LAFL
 Sbjct: 61 AVAVKOLLNKLTFEGIKNDLNEIDGHRVHVGSEKFSDSVLVDTETIKEDISELAFL 120
 30 Query: 121 HNPAAAGIIRAFREILPDITSCVCFDTSFHTSMKHTYLYPIPKQYTYDVKVKYGAHGT 180
 HNP GI+AF++LE++ +V VEDT+FH +M + +YLY +P +YY + +RKY HET
 Sbjct: 121 HNPANIVGIIKAFKVELNVPFAVAVDIAFHQTMPEQSYLSLPTEYFEKGIKRYGPHGT 180
 Query: 181 SHKYVAQEAARKLGRPLEELKLITAHNGVSVITANYHGKSVDTSMGFTPLAGVMMSTRS 240
 SHKYV + AA++LGRPL+L+LI+ H+NG SI A GKS+DTSMGFTPLAG MSTRS
 35 Sbjct: 181 SHKYVTERAAELGRPLKDLRLISCHLNGASIAAEBGKSIDTSMGFTPLAGVMMSTRS 240
 Query: 241 GDIDPAIIPYLIQDPELKDAADVNNLNKSKSLGSGVSGISDMRDIEAGLQEDNPDAVL 300
 G+IDPA+IPY++E+ + D +V+N LNKSKSL G+SG SSD+RDI +E N A
 Sbjct: 241 GNIDPALIPYINKEGTQAD--EVNTLNKSKSLGISGFSDDLRIEATKCGNERAET 300
 40 Query: 301 AYNIFIDRIKKICQYFAVINGADALVFTAGMGENAPLMQDVIGLWFGMDIDPE-IN 359
 A +F RI K IG Y A +G DA++FTAG+GEN+ +R+ V+ GL +G+ DP N
 Sbjct: 299 ALEVFASRIHKYIGSYAARMGVDAIIFTAGIGENSEVEVRERVLRLFMGVYWDPALIN 358
 45 Query: 360 VFGYGRDITSPKSKVKVLVIPTDEELMIARDVRL 394
 V G IS P S VKV+I TDE+ IARDV RL
 Sbjct: 359 VRGEAFISYPHSPVKVMIIPTDEEVMIAADVRL 393

An alignment of the GAS and GBS proteins is shown below:

Identities = 332/395 (84%), Positives = 365/395 (92%)
 50 Query: 1 MSKTIAINAGSSSLKWLQYMPPEAVLAQGIIRIGLAKDSISTVKYDGKKSEQLIDIH 60
 MSKTIAINAGSSSLKWLQY+MPPE VAA+GIIRIGLAKDSISTVK+D KK+EQI+DI DH
 Sbjct: 1 MSKTIAINAGSSSLKWLQYMPPEAVLAQGIIRIGLAKDSISTVKYDGKKSEQLIDIH 60
 55 Query: 61 TQAVKILLNDLHFGIIAAYDEITGVGHRVAGGELFKESVVDKVLQIEELSLVLAFL 120
 T+AVKILL DL GII ++EITGVGHRVAGGE FKES +V+DKV+Q+BELS LAFL
 Sbjct: 61 TEAVKILLNDLHFGIIAAYDEITGVGHRVAGGELFKESVVDKVLQIEELSLVLAFL 120
 60 Query: 121 HNPAAAGIIRAFREILPDITSCVCFDTSFHTSMKHTYLYPIPKQYTYDVKVKYGAHGT 180
 HNP AAAGIIRAF+ILPDITSCVCFD+FT+M HTYLYPIPKQYTYDVKVKYGAHGT
 Sbjct: 121 HNPAAAGIIRAFREILPDITSCVCFDTSFHTSMKHTYLYPIPKQYTYDVKVKYGAHGT 180
 Query: 181 SHKYVAQEAARKLGRPLEELKLITAHNGVSVITANYHGKSVDTSMGFTPLAGVMMSTRS 240

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SH+YVAQSAAK LGRPLEELKLITAH+GNGVSVITANYHG+S+DTMGFTPLAGFMQSTRS
 Sbjct: 181 SHKYVAQSAAKMLGRPLEELKLITAHIGNVSVITANYHGKSVITMGFTPLAGFMQSTRS 240

Query: 241 GDIDFAIIPYLAVNDPRLDAAAVVMMLNKQSLGVSSTSSMDRIAGLQSKDPNAV 300
 GDIDFAIIPYL+ DPEL+DAA VVMMLNK+SLGVSG SMDMDRIAGLQ +P+AVL
 Sbjct: 241 GDIDFAIIPYLIEQDFELKDAAVVMMLNKKSLSGVSGISSMDMDRIAGLQKINPNAV 300

Query: 301 AYNVFDIRIKKFIQQYFAVANGADAIIFTAGMGENAPLMRQDVLAGLSWFGIELDPEKIV 360
 AYN+FDIRIKK IQQY AVANGADA++PTAGMGENAPLMRQDVI GL+WFG++DPEKIV
 Sbjct: 301 AYNVFDIRIKKCIQQYFAVANGADALVFTAGMGENAPLMRQDVIGSLTWFGMDIDPEKIV 360

Query: 361 FGYPGDITKPSQKVKVLVITDEELCIARDVERLK 395
 FG Y GDI+ P+SKVKVLVI DEEL IARDVERLK
 Sbjct: 361 FGTRGDISTPESKVKVLVISTDEELCIARDVERLK 395

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 121

A DNA sequence (GBSx0126) was identified in *S.agalactiae* <SEQ ID 409> which encodes the amino acid sequence <SEQ ID 410>. This protein is predicted to be repressor protein. Analysis of this protein sequence reveals the following:

Possible site: 17
 >>> Seems to have an uncleavable N-term signal seq

----- Final Results -----
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CA849550 GB:AJ248284 repressor protein, putative [Pyrococcus
 abyssi]
 Identities = 39/64 (60%), Positives = 49/64 (75%)

Query: 1 MKNLSQKLRKRLSQAEALVALGVTRQTIIISLEKKYTSLELAFKIRYFDKQIEEVF 60
 MKN L+ +R+ L+Q ELA LGVTRQTII++EK KY SL LAFKIR+ F +IE++F
 Sbjct: 1 MKNRLREFREKYGLTQELARILGVTRQTIIAIEKGKYDPSLELAFKIRFFGVRIEDIF 60

Query: 61 IYTE 64
 IY E
 Sbjct: 61 IYEE 64

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 411> which encodes the amino acid sequence <SEQ ID 412>. Analysis of this protein sequence reveals the following:

Possible site: 40
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.4344 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 29/66 (43%), Positives = 44/66 (65%)

Query: 1 MKNLSQKLRKRLSQAEALVALGVTRQTIIISLEKKYTSLELAFKIRYFDKQIEEVF 60
 +KN L+LR ++Q R+R GV+RQTI +E+ +YT S+ +A KIA+ F + +EEVF
 Sbjct: 10 LQNRLKELRARDGINQETMAKLAGVSRQTISLIERNEYTPSVIIAMKIAKVFQFVEEVF 69

Query: 61 IYTRSE 66
 E E
 Sbjct: 70 RLVEVE 75

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 122

A DNA sequence (GBSx0127) was identified in *S.agalactiae* <SEQ ID 413> which encodes the amino acid sequence <SEQ ID 414>. Analysis of this protein sequence reveals the following:

```
Possible site: 32
>>> Seems to have an uncleavable N-term signal seq
  INTEGRAL    Likelihood = -8.97    Transmembrane    45 - 61 ( 41 - 66)
  INTEGRAL    Likelihood = -8.65    Transmembrane    14 - 30 ( 11 - 37)
  INTEGRAL    Likelihood = -7.80    Transmembrane   123 - 139 ( 118 - 145)
  INTEGRAL    Likelihood = -3.24    Transmembrane   177 - 193 ( 177 - 194)
  INTEGRAL    Likelihood = -0.85    Transmembrane    81 - 97 ( 81 - 97)

----- Final Results -----
bacterial membrane --- Certainty=0.4588 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
```

A related GBS nucleic acid sequence <SEQ ID 9491> which encodes amino acid sequence <SEQ ID 9492> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```
>GP:BA11325 GB:D78257 ORF8 [Enterococcus faecalis]
Identities = 48/120 (40%), Positives = 69/120 (57%), Gaps = 5/120 (4%)

Query: 104 MQGVKDTANQTVIMELTKQLPLALMLIPAIIGAPIMEEIIIPRYIIPKRIIPAKHQKQWPI 163
      MQG TAN + +++L + L+++ I AFIMEEII+FR I L + +I
Sbjct: 1 MQGHTTFANDSTLILKLPQGVSPVLVLLGLIAAPIMEEIVFRGGIIGYIVENNALLAILI 60

Query: 164 GTLAFALIHSPSDIGSPFIYAGMGAILSPVYYKTHELVYSIMHFINN-----ALAYSVL 218
      + F +IH P++ SF +Y MG ILS YKTI+ L SI IHP+NN AAY ++
Sbjct: 61 SSFLFGIIGHPTNFISPGMYFPNGIILSVSYKTKOLRVISIHFLNLPALAIAYGLI 120
```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 415> which encodes the amino acid sequence <SEQ ID 416>. Analysis of this protein sequence reveals the following:

```
Possible site: 24
>>> Seems to have an uncleavable N-term signal seq
  INTEGRAL    Likelihood = -11.41    Transmembrane    12 - 28 ( 1 - 30)
  INTEGRAL    Likelihood = -9.98    Transmembrane    41 - 57 ( 33 - 64)
  INTEGRAL    Likelihood = -8.33    Transmembrane   128 - 144 ( 121 - 151)
  INTEGRAL    Likelihood = -7.96    Transmembrane    83 - 99 ( 76 - 103)
  INTEGRAL    Likelihood = -3.77    Transmembrane   208 - 224 ( 207 - 230)
  INTEGRAL    Likelihood = -2.13    Transmembrane   182 - 198 ( 182 - 199)

----- Final Results -----
bacterial membrane --- Certainty=0.5564 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
```

The protein has homology with the following sequences in the databases:

```
>GP:BA11325 GB:D78257 ORF8 [Enterococcus faecalis]
Identities = 47/120 (39%), Positives = 70/120 (58%), Gaps = 8/120 (6%)
```

Query: 105 GQVSDANDAIHTLARLIKGGFPLYTALFVLVIAFIAPIMEELVFRGFPMDLFEKGSILK 164
 G +AND+ TL +L G P+ L VL++ APIMEE+VFRG + L+ +L
 Sbjct: 3 GHTTTANDS---TLKLFSGVSPV---LVLLGLIAAPIMEEIVFRGSIIGYLVENNAL- 55

5 Query: 165 VAGLVTSLVFALPHA-TNSVEFIMYSCMGIFLVFVAYQRRGNLKDAILLHIFNNLEIVILL 223
 +A L++S +F + H TN + F MY MGI L V+Y + +L+ +I +H NNL I +
 Sbjct: 56 LAILISSFLPGIIGPINFISFGMYFPFGIILSVSYTKDKLEVSIHFLNNLPFAIAI 115

An alignment of the GAS and GBS proteins is shown below:

10 Identities = 72/229 (31%), Positives = 114/229 (49%), Gaps = 24/229 (10%)

Query: 11 KGKILALLIAFLVINGLV-PILAVLLKHNYQFPPTSILLIGL-----ELLIATLFLY 62
 KG I L IA L+I +V +L+ LL+ + P IG+ +LI+ LW
 Sbjct: 2 KGFINVLKIAVLILAMVFNVLDMILLQKHDI PWLWNGSIGIFYLTVGSGVLIVAGLY 61

15 Query: 63 YAKVQIIRWKALLTRKALVT---ILGWLSLEVQIIGYLIMTM-QGVKDTANQTVMIE 118
 AK I+ + + LW + L WL +RV I+G L+ + G +AN I
 Sbjct: 62 QAKQDTPIKQKIM----RLVDWGIALFWLIRVIAIVGTLVNLQWSSQVSDANDAIHT 117

20 Query: 119 LTKQL----PLALMLFAIIG--APIMEEIIPKYIIPKELF-AKHQKGVIGTLAFALI 171
 L + + PL L +I APIMEE+FR +LF K K + +L EAL
 Sbjct: 118 LARLIKGGFPLYTALFVLVIAFIAPIMEELVFRGFPMDLFEKGSILKVLAVTSLVFALP 177

25 Query: 172 HSPSDIGSFLIYAGGAILSPVYVYKTEHLYSIIHFINNALYSVLIS 220
 H+ + + FI+Y+ MG L Y + +L+ +I+H NN + +L+S
 Sbjct: 176 HATNSV-EFIMYSCMGIFLVFVAYQRRGNLKDAILLHIFNNLEIVILLAS 225

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

30 Example 123

A DNA sequence (GBSx0128) was identified in *S.galactiae* <SEQ ID 417> which encodes the amino acid sequence <SEQ ID 418>. Analysis of this protein sequence reveals the following:

Possible site: 14
 >>> Seems to have no N-terminal signal sequence

35 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.0826 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAC06504 GB:AB000676 pyrroline carboxylate reductase [Aquifex
 seolicus]
 Identities = 97/259 (37%), Positives = 159/259 (60%), Gaps = 4/259 (1%)

45 Query: 1 MKIGIIGVGRM--ASAIQQLWOTONDIIISGSLERSKRIARLDVTVYASHOSLINDA 58
 M++GI+G G M A A+ K + +II++ E+ +A + + +A + L + +
 Sbjct: 8 MRVIGVPGMGQAFALCFSKLKGKNIIVIDKQEK-RNLATMGIAFASDVKFLADNS 66

50 Query: 59 DIIMLGIKPLFEKVLPLDITKPII-SMAGISLARISQLTRSDLPILRIMENNAQIL 117
 D++++ +KP+ +VL L K II S AG+S+ ++ D +R+MEN+N +
 Sbjct: 67 DVVLVAVKPKDSQVVLQKLDYKGIILSMAGVSIERMEKILGDKICIVRVMVNVAVG 126

55 Query: 118 QSCTAICYNHNSVDELRLAKEITDSFGSSPDIAETNFTPTMALGSSPFIYIPLTALA 177
 AI N++S+E R +E+ S G+ + I E ED FTALAGS PA+++ PI+ALA
 Sbjct: 127 SGVVAITDNGNLSEERSKVEELLISOSTLYRIKERLFDATLAGSGPAFVSPIDALA 186

Query: 178 KAGVKYGFPEQALSIVGTVLASSONLLQQONSTSLYDNICSPGGTITAGLLDLERNG 237
 AGV GP EQNL I TV+ S++ L + Q + +LI + SPGGTIT G+ LE+ G
 Sbjct: 187 LAGVHQGPSVEQALRIALDTVMGSAKLLKEPQVNEELIAKVTSFGSTTIBGKIYDEKG 246

-201-

Query: 238 LTHSVISADATIEKAKKL 256
 +V+ I+ T+ KAKKL
 Sbjct: 247 FKGTVMBCINRTSQKAKL 265

- 5 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 419> which encodes the amino acid sequence <SEQ ID 420>. Analysis of this protein sequence reveals the following:

Possible site: 50
 >>> Seems to have no N-terminal signal sequence

- 10 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1043 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

- 15 An alignment of the GAS and GBS proteins is shown below:

Identities = 180/256 (70%), Positives = 208/256 (80%)

- Query: 1 MKIGIIGVGKMASAIIGLKTQHDIIISGSLERSKEIAERLDVTAESHQSLINQADI 60
 20 MKIGIIGVGKMASAIIGLKTQ H++IISGS LERSKEIAE+L +YA SHQ LI+Q D+
 Sbjct: 1 MKIGIIGVGKMASAIIGLKTQPHLIIISGSLERSKEIAEQALPYAMSHQDLIDQVLD 60
 Query: 61 IMLGIKPOLPEKVLPLDITKPIISMAGISLARISQLTRSDLPRLIRIMFNINAQILQSC 120
 ++LGIKPOLPE VL FL +PIISMAGISL RL+ DLPL+RIMFN+NAQILQSC
 25 Sbjct: 61 VILGIKPOLFETVLKPLHFKQPIISMAGISLARISQLTRSDLPRLIRIMFNINAQILQSS 120
 Query: 121 TAICNNHVSDELRLQAKELTDSFGSSFDIAETNFDTPALAGSSPAYIYLFIEALAKAG 180
 TA+ N VS EL+ ++TDSFGS+FDI+E +FDTPTALAGSSPAYIYLFIEALAKAG
 30 Sbjct: 121 TALTGNALVQELQARVRLTDSFGSTFDISEKDFDTPTALAGSSPAYIYLFIEALAKAG 180
 Query: 181 VKYGPKEQALSIQVGTQVLASSNQLGQNSDLDINICSPGGTTIAGLLDLEKNGLTH 240
 VK G PK +AL IV QTVLAS+ NL S D ID ICSPGGTTIAGL++LE+ GLT
 35 Sbjct: 181 VKKGIKPAKALEIVTQTVLASASNLKTSQSPHDFIDAICSPGGTTIAGLLERLGLTA 240
 Query: 241 SVISADATIEKAKKL 256
 +V SAID TI+KAK L
 35 Sbjct: 241 TVSSAIDKTIIDKAKSL 256

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

40 Example 124

A DNA sequence (GBSx0129) was identified in *S.agalactiae* <SEQ ID 421> which encodes the amino acid sequence <SEQ ID 422>. Analysis of this protein sequence reveals the following:

Possible site: 58
 >>> Seems to have no N-terminal signal sequence

- 45 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3405 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

50

The protein has homology with the following sequences in the GENPEPT database:

>GP:CPA56994 GB:X81089 glutamyl-aminopeptidase [Lactococcus lactis]
 Identities = 219/354 (61%), Positives = 273/354 (76%), Gaps = 1/354 (0%)
 55 Query: 3 DLPNKKITVTELDGLAGVHNHINFLRQETLPLVDQVETDGLGGI PGVKNTHETNAPKVM 62
 +LF+K+K +TE+ +G+R +R++L+ +L Q E IGLGTF K+ NAP++M
 Sbjct: 2 ELFDKVKALTEIQATISGFBGPFVRDYLAARMVELQYQPEFGLOGIFVTKASKVENAPRIM 61
 Query: 63 VAAHMDVGVFVSHIQPDGTFRVLEVGGMNPLVSSQRPFLYTRSGDAIPVISGSPVPHF 122

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- VAAHMEVGFPMVS I+ DGTFRV+ +GWNPLVVS QRFTL+TR+G IPV+G +P+H
 Sbjct: 62 VAAHMEVGFPMVSSIKADGTFRVVPLGWNPLVVSQRFTLPTRTGKKIPVVTGGLP+H 121
- Query: 123 LRGGSGGTTLPKISDIFVFDGGFTDKNRAESFGLAPGDIIIVPKSFTLLTANQKHIMSKAWD 182
 LRG +P ISDIF+FDG F+ EA FGLA GD+I+P+ETTL+AN K+I+SKAWD
 Sbjct: 122 LRGTGVTQIPATISDIFDGAFAENAAEAFFGLAQGLDIIIPETETLLSANGKNIISKAWD 181
- Query: 183 NRYGVLMVTELLKSLKQSLNTLITAGANVQREVGLRGAVHSTTKFNPDIPLAVDCSPA 242
 NRYG LM+ EL+ L D+ L TLL GANVQREVGLRGA VSTTKNPD+F AVDCSPA
 Sbjct: 182 NRYGCLMLELLEFLADKELFVTLITAGANVQREVGLRGAVSTTKFNPDLFFAVDCSPA 241
- Query: 243 DIYG-EQGGIGSGTILRFYDPGHIMLKMNRDPLLTAREAGIKYQYYAANGSTDAAGAH 301
 D +G +G+GGGT +RF+DPGHIML M++FL TA A +K Q Y A GGTDAAGAH
 Sbjct: 242 DTFGDNGRLGSGTILRFDPGHIMLQMKNFLLDTAMHAKVKTQYVMNKGSTDAAGAH 301
- Query: 302 KNSOIPSTTIGVCARYIHSHQTLVAMDDFLQAQAYLQAINVKLDRSTVDIIKGY 355
 N G+PSTTIGV ARYIHSHQT++ +DDFLQAQ +L+AI+ L+ V IK Y
 Sbjct: 302 ANGGVPSTTIGVARYIHSHQTLFNHDDFLQAQTFLRALITSLMTKVAEKNY 355
- A related DNA sequence was identified in *S.pyogenes* <SEQ ID 423> which encodes the amino acid
 sequence <SEQ ID 424>. Analysis of this protein sequence reveals the following:
- Possible site: 55
 >>> Seems to have no N-terminal signal sequence
- Final Results -----
 bacterial cytoplasm --- Certainty=0.2747 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
- An alignment of the GAS and GBS proteins is shown below:
- Identities = 276/355 (77%), Positives = 322/355 (89%)
- Query: 1 MSDLEFNKIKTVTELLGDIAGYEHNIENFLRQBITPLAVDQVETDGLGIFGVNTHETNAPK 60
 M+DLF+KIK VTELLGDIAGYEH++R+LR +ITPLVD+VETDGLGIGF++ AP+
 Sbjct: 1 MIDLFSEKIKVTELLGDIAGYRHSVRDYLETKITPLAVDEVETDGLGIFGRDSKARKAPR 60
- Query: 61 VVAAHMEVGFPMVSHIQPDGTFRVLEVGGWNLVVSQRFTLYTRSGDAIPVISGSVPP 120
 ++VAAHMEVGFPMVS I+ DGT RV+ +GWNPLVVSQRFTL+TR+G IP+IGSGVPP
 Sbjct: 61 ILVAAHMEVGFPMVSDIKVGLTRVVG+GWNPLVVSQRFTLYTRTQVPIILISGSVPP 120
- Query: 121 HFRLGGSGGTTLPKISDIFVFDGGFTDKNRAESFGLAPGDIIIVPKSFTLLTANQKHIMSKA 180
 HFRLG +G +LP I DIVFDGGFTDK NAE FGI PGDII+P+SETLLTANQK+I+SKA
 Sbjct: 121 HFRLGANGSASLPHIEDIVFDGGFTDKAASERFGITPGDIIIPQSTLLTANQKNIISKA 180
- Query: 181 WENRYGVLMVTELLKSLKQSLNTLITAGANVQREVGLRGAVHSTTKFNPDIPLAVDCSP 240
 WENRYGVLM+TE+L+LK Q L+NTLITAGANVQREVGLRGAVHSTTKF+P++F AVDCSP
 Sbjct: 181 WENRYGVLMITEMLRLEGGQDLNLTITAGANVQREVGLRGAVHSTTKFDPKPLFFAVDCSP 240
- Query: 241 AGDIYGEQGGIGSGTILRFYDPGHIMLKMNRDPLLTAREAGIKYQYYAANGSTDAAGAH 300
 AGDIY G IG+GTL+RFYDPGH+MLKMNRDPLLTAREAG+ +QYY CGTDAAGAH
 Sbjct: 241 AGDIYGNPGTIGDGTILRFYDPGHVMIKMNRPFLITAREAGVNFQYYCGGTDAGAH 300
- Query: 301 LKNSGIPSTTIGVCARYIHSHQTLVAMDDFLQAQAYLQAINVKLDRSTVDIIKGY 355
 L+N G+PSTTIGVCARYIHSHQTLVAMDDP++AQA+LQAI+ KLDRTVD+IK Y
 Sbjct: 301 LQNGVPSTTIGVCARYIHSHQTLVAMDDPFAQAFLQAIKKLDRSTVDIIKCY 355

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 125

- 60 A DNA sequence (GBSx0130) was identified in *S.galactiae* <SEQ ID 425> which encodes the amino acid sequence <SEQ ID 426>. Analysis of this protein sequence reveals the following:

Possible site: 26
 >>> Seems to have no N-terminal signal sequence

- 5 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1672 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

- 10 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 126

- 15 A DNA sequence (GBSx0131) was identified in *S.agalactiae* <SEQ ID 427> which encodes the amino acid sequence <SEQ ID 428>. Analysis of this protein sequence reveals the following:

Possible site: 31
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -2.28 Transmembrane 18 - 34 (17 - 34)

- 20 ----- Final Results -----
 bacterial membrane --- Certainty=0.1914 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

- 25 The protein has no significant homology with any sequences in the GENPEPT database.

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 429> which encodes the amino acid sequence <SEQ ID 430>. Analysis of this protein sequence reveals the following:

- 30 Possible site: 21
 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -6.16 Transmembrane 12 - 28 (8 - 30)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.3463 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

An alignment of the GAS and GBS proteins is shown below:

- 40 Identities = 30/91 (32%), Positives = 48/91 (51%)
 Query: 13 MKNKKILPGTGLAVGGLAAGYTLTKKVTDYKRQQITQTTLRPFSSQMGDIQVFYFNEPE 72
 M KKI +G+ G L G + D +R+Q+T+ LR PFS +G I+V Y N +
 Sbjct: 4 MSKKKIGMISGIPGSLATGLGIVIKDYQDQRQRQTRDLRTFPSPILQIEVLVYNPCQ 63
 Query: 73 SDIKMTSGGLVLEGRIFEFYIRQGVLDYVE 103
 SGG+V+ +G+ ++F Y + + E
 Sbjct: 64 VRQDYISGGVMSGGKQYQFTYHSRQISPER 94

- 50 A related GBS gene <SEQ ID 8497> and protein <SEQ ID 8498> were also identified. Analysis of this protein sequence reveals the following:

Lipop Possible site: -1 Crend: 4
 SRCFLG: 0
 MoG: Length of UR: 21

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```

Peak Value of UR: 2.30
Net Charge of CR: 3
McG: Discrim Score: 6.28
GVH: Signal Score (-7.5): -1.46
5   Possible site: 19
    >>> Seems to have a cleavable N-term signal seq.
    Amino Acid Composition: calculated from 20
    ALOM program count: 0 value: 22.60 threshold: 0.0
    PERIPHERAL Likelihood = 22.60 29
10   modified ALOM score: -5.02

*** Reasoning Step: 3

Rule gp01
15   ----- Final Results -----
        bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
        bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
20   bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

SEQ ID 8498 (GBS214) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 40 (lane 3; MW 13.9kDa). It was also expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 46 (lane 6; MW 39kDa).

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 127

A DNA sequence (GBSx0132) was identified in *S.agalactiae* <SEQ ID 431> which encodes the amino acid sequence <SEQ ID 432>. This protein is predicted to be thioredoxin H1 (trxA). Analysis of this protein sequence reveals the following:

```

30   Possible site: 40
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
        bacterial cytoplasm --- Certainty=0.2350 (Affirmative) < succ>
35   bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
        bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

40   >GP:BA006972 GB:AP001518 thioredoxin H1 [Bacillus halodurans]
    Identities = 47/90 (52%), Positives = 66/90 (73%)

    Query: 14 IDSTKKVVFPTADWCPDCQFLYPVPMSEIEKDFSPFVFRVNRDDYIELAQCNILGIPS 73
    + + + VVF P+AWCPDC+ I P +P +E+ + + F VNRDD+IEL Q+ +IPGIPS
    Sbjct: 13 VKNQENVVFLPSADWCPDCRVIRPFLPELEQTYDEYQFYVNRDDFIELQELDIPGIPS 72

    Query: 74 FVVVENGQELGRVKNRKNRKTAEITKFLAE 103
    F+ NG+E R V+K+RKTK EI +PL E
    Sbjct: 73 FLFYENGGERSRFVSKDRKTKKEIRFLTE 102

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 433> which encodes the amino acid sequence <SEQ ID 434>. Analysis of this protein sequence reveals the following:

```

    Possible site: 35
    >>> Seems to have no N-terminal signal sequence

55   ----- Final Results -----
        bacterial cytoplasm --- Certainty=0.1997 (Affirmative) < succ>
        bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

```

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 70/102 (68%), Positives = 81/102 (78%)

Query: 1 MILPESGYEELAAVIDSTKKVFFPTADWCDDQPIYVPMPSIEKDFSDVFPVVRNDDVI 60
MI P SYR 4A I+ K+V FFTADWCDDQPIYV+MP IR + +D FV VNED +I
Sbjct: 1 MIRPTSYESLATLIKEDKLVLFPTADWCDDQPIYVIMDEIRAELEMTFTVCVNRDQFI 60

Query: 61 ELAQQNNIFGIPSPVVEGQELGRLVNRKTKARITKFLA 102
E+AQ+WNIFGIPSPV+E GQE+GRLVNRK RTK EI FLA
Sbjct: 61 EVAQNNIFGIPSPVVEKGQGVRLVNRKTKTEIMHFLA 102

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 128

A DNA sequence (GBSx0133) was identified in *S. agalactiae* <SEQ ID 435> which encodes the amino acid sequence <SEQ ID 436>. This protein is predicted to be phenylalanyl-tRNA synthetase beta subunit, non-spirochete. Analysis of this protein sequence reveals the following:

Possible site: 47
>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1310 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AA000291 GB:AF008220 YtpR [Bacillus subtilis]

Identities = 78/196 (39%), Positives = 125/196 (62%), Gaps = 1/196 (0%)

Query: 5 YNREHVGDITLMVIVKDSQAKLDVDRGQVAVRYLQDSKETVAMNIFVSSLIIVIEGAGQ 64
YN+E VGDITL++ ++D +L ++ G V +++ ++KET +NIF SS + I+ G
Sbjct: 5 YNREHVGDITLLISLQDVIREQLGYEKHGQVVKIPNNETKETTGFNINASSYLITIDENGP 64

Query: 65 ITLSQDQIKILNABELKEGFEDSLVNNIEPTFVVAQIKETIDHPDSHRLHQAQINDGK 124
+LS+ ++ +N L + G E++LV ++ P FVV ++ HF++D L +C+ + + +
Sbjct: 65 VALSETFVQDVNELLNRRNGVEETLVDLSPKFVGVGVSEKHKFNADKLSVCKVAVNGE-E 123

Query: 125 TVQIVOGAPNAPSVGLKTVAAALPGAMVPNGSLIFPGKLRGDSFGMLCSARRLALPNAPQV 184
T+QIVOGAPN G K V A GA+MP+G +I +LRG S GM+CSA+EL LP+AP
Sbjct: 124 TLQIVOGAPNVDQQGVVAVRVGAVMPSGLVTKDAELRGVPSGSMCSA+GLDLPDAPAE 183

Query: 185 RGIIELSDQVIGESF 200
+GI+ L G++F
Sbjct: 184 KGIIELSGDYRAGDAF 199

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 437> which encodes the amino acid sequence <SEQ ID 438>. Analysis of this protein sequence reveals the following:

Possible site: 47
>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood = -1.49 Transmembrane 90 - 106 (90 - 107)

----- Final Results -----

bacterial membrane --- Certainty=0.1595 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP:BAB06970 GB:AP001518 phenylalanyl-tRNA synthetase (beta subunit)
[Bacillus halodurans]
Identities = 84/196 (42%), Positives = 124/196 (62%), Gaps = 1/196 (0%)

5 Query: 5 YNKEQVGDVLMVILQDTKDIKQVERKGKVARVFAESGKTLAWNIPASSLITIEGNOQ 64
YN++GD++++++RER+GV R++ +GKT +N+P AS GG
Sbjct: 5 YNERGIGDTLILVIDEVEPANRAYERQSDVVRIYHLSGTGKTGYNLPHASKYGFEPNGQL 64

10 Query: 65 IFLIDENLARLNALAKJKEGFSERLEPIGVPPVVGQIVEMVAHPDSHNLNQVAIGEDQ 124
+LTD +A L K G + LE + P FVVG + HP++D L+I+V +G D
Sbjct: 65 LRLTDSLVAITLQAFQKNGVNNWILEVDLSPKFVVGFGVQSKDKHPNADKLSTCKVDVGSQ 123

15 Query: 125 TVQIVAGAPNAALGLKTIIVALEGAIMPNGLIFPGKLRGRRSYGMNCSPRRLALPNAPQK 184
T+QIV GAPN G K +VAL GA+MP+G +I P LRG S GM+CS +ELALP+AP++
Sbjct: 124 TLQIVOGAPNVEAGQKVVALEGAVMPSGLWIKPTSLRGVSGTGMCSAKRLALPDAPES 183

Query: 185 RGIIEFDESAVVGEAF 200
+GI+ D+S VG +F
20 Sbjct: 184 KGIIVLEDSYEVGTSF 199

An alignment of the GAS and GBS proteins is shown below:

Identities = 133/207 (64%), Positives = 167/207 (80%)

25 Query: 1 MIPTYNRERVGDITLWIVKDSQGAKLVDPRGQVARVYLQDSKETVAMNIEFVSSLIVIE 60
MIF YN+E VGD LMVI++D++ K V+R+G+VARV+ ++S +T+AMNIEF SSLI IE
Sbjct: 1 MIFAYNKEQVGDVLMVILQDTKDIKQVERKGKVARVFAESGKTLAWNIPASSLITIE 60

30 Query: 61 GAGQITLSDQDIKILNARLLKJEGFEDSLVNNIEPTVVAQIKKIIIDHPDSHNLCOAEI 120
G GQI L+D+++ LNARL KEQF + L + P FVV QI E++ HPDSHNL+ICQ I
Sbjct: 61 GNGQITLIDENLARLNALAKJKEGFSERLEPIGVPPVVGQIVEMVAHPDSHNLNQVAI 120

Query: 121 NDGKTVQIVOGAPNAGVLKTVAALEGAMMPNGSLIFPGKLRGDSFGMLCSARELALFN 180
+ +TVQIV GAPNA++GLKT+ ALPGA+MPNGSLIFPGKLRGE+S+GM+CS RELALFN
35 Sbjct: 121 GEDQTVQIVOGAPNAGVLKTVAALEGAIMPNGLIFPGKLRGEESGMNCSPRELALFN 180

Query: 181 APQVRGIIELSDQVIVGESFDDANKHWK 207
APQ RGIIE + +VGE+FD KHWK
40 Sbjct: 181 APQVRGIIEFDESAVVGEAFDEAKHWK 207

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 129

A DNA sequence (GBSx0135) was identified in *S.galactiae* <SEQ ID 439> which encodes the amino acid sequence <SEQ ID 440>. Analysis of this protein sequence reveals the following:

Possible site: 30
>>> Seems to have no N-terminal signal sequence

---- Final Results ----

50 bacterial cytoplasm --- Certainty=0.3052 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:BA081904 GB:U92974 unknown [Lactococcus lactis]
Identities = 69/241 (28%), Positives = 117/241 (47%), Gaps = 15/241 (6%)

55 Query: 7 YKENLAKPWGKIQYKTFAGL--SHIKNQNVLDGAGFCLTECHLAKEN-NVTALRPNPK 63
Y E+ KIWG++ Y++ F QL + K+ +L FG+GF TE L++ VT EP+ +
60 Sbjct: 23 YAEVFEKPGWRMFYDILFLPQLILPNTKDSKILSFGSGRGRTYTLFEEQGFVETGYDVE 62

-207-

Query: 64 LLYDNQSDNIYKILASVEALRD-LPQQSFITIIICNVLEVIDKINHNPAYDFEFSRLIKFN 122
 L ++ G+++ + + + +D I+ HNVLEF+ + + L+
 5 Sbjct: 83 KLEHMSDQTFRQLTGTPDDFAETVQNERIVLILHNVLEFV--IDRKVLELLLELLTDG 140

Query: 123 GELSLIKHNTIKILQSVIPSNDSAMELLINKEANFKASFDQGNIT-----LEELNQ 177
 G LS++KH+ G +++ ++ A+++ EA AS + G+I L +
 10 Sbjct: 141 GTLSIVKHSKYGSMTEMAAGRDNPQALDVEYNERA---VASINHGSDILVYDDDLWLTDFVA 197

Query: 178 NINLLVRYQGHITFYSYSLQW-HFKTETGWLAKMLAELSVADKAPYKDIAFLQHTLKKS 237
 N L ++ GIR FZ + N K W ML +E VA +A L H+ KGS
 10 Sbjct: 198 NYKLKLBKGGRHIFTGISQNAIKETENWYQFMLEKQKVAKDCTLYPVARLHLLFKKS 258

No corresponding DNA sequence was identified in *S.pyogenes*.

- 15 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 130

A DNA sequence (GBSx0136) was identified in *S.agalactiae* <SEQ ID 441> which encodes the amino acid sequence <SEQ ID 442>. Analysis of this protein sequence reveals the following:

20 Possible site: 58
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 25 bacterial cytoplasm --- Certainty=0.3479(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

30 >GP:AAF74079 GB:AF212845 putative single stranded binding protein
 [Lactococcus lactis bacteriophage ul36]
 Identities = 64/141 (45%), Positives = 92/141 (64%), Gaps = 10/141 (7%)

Query: 1 MYNKVIMIGRLTAKFEMVKIPTDKSVTRATVAVNRFRFGSGNGERRADFINVVMGRLAET 60
 M N V ++GR+T +PE+ TP +K+V T+AVNR FK +NGRREADFI+ V+WG+ AE
 35 Sbjct: 1 MINNVITLVGRITKPELRYTPQNKAVALFTLAVNRAPFNANGERRADFTSCVWGKSAEN 60

Query: 61 LASYGTKSLISIDGELATRIKYE-KDGOTHYITEVLASSPOLLESRAQ-----RAM 110
 LA++ KG LI + G ++TR YE + GQ YITEV+AS+PQ+LE C +
 40 Sbjct: 61 LANWTHKGQLIGVIGNICTRYENCCQQRVYITEVVASNPQVLEKSNANGERISNPASK 120

Query: 111 RENNVSDLSLDLVEEELPF 131
 +NN S + + ++LPP
 Sbjct: 121 PQNDSFGSDPMEISDDLPF 141

- 45 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 443> which encodes the amino acid sequence <SEQ ID 444>. Analysis of this protein sequence reveals the following:

Possible site: 32
 >>> Seems to have no N-terminal signal sequence

50 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1817(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

- 55 An alignment of the GAS and GBS proteins is shown below:

Identities = 102/131 (77%), Positives = 116/131 (89%)

Query: 1 MYNKVIMIGRLTAKFEMVKIPTDKSVTRATVAVNRFRFGSGNGERRADFINVVMGRLAET 60

-208-

MYNKVI IGRLL AKPB+VKT TDK V R ++AVNRFPK ++GERADPT+VV+WG+LAET
 Sbjct: 1 MYNKVIAIGRLVAKPBLVKTATDKHVARLSLAVNRFPKASGERADPTISVVVGKLAET 60
 Query: 61 LASYCTGKSLSIDGELRTRKYEDQCTHYITEVLASSFQLLESRAQRAMRNNVSDLS 120
 L SY +KGLS+SIDGELRTRKY+KDGQ HY+TEVL SFQLLESRAQRAMRNNV+ DL
 Sbjct: 61 LVSYASKGSLMSIDGELRTRKYDKGQHYHYITEVLCSFQLLESRAQRAMRNNVNTDLV 120
 Query: 121 DLVLEETLPP 131
 DLVLEA+ LPP
 Sbjct: 121 DLVLEETLPP 131

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 131

- 15 A DNA sequence (GBSx0137) was identified in *S. agalactiae* <SEQ ID 445> which encodes the amino acid sequence <SEQ ID 446>. Analysis of this protein sequence reveals the following:

Possible site: 49
 >>> Seems to have no N-terminal signal sequence

- 20 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.2235 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

- 25 A related GBS nucleic acid sequence <SEQ ID 9493> which encodes amino acid sequence <SEQ ID 9494> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

- >GP:CACL3072 GB:AI445503 putative hydrolase [Streptomyces
 coslicolor]
 30 Identities = 63/179 (35%), Positives = 91/179 (50%), Gaps = 2/179 (1%)
 Query: 33 IIFMDGVIVDSSEYTFLNKTEMLREBGI-DTDSYQYQYKITTPEFMQAMKEEGLPK 91
 +IFD+EG +VDSH ++ L E G+ D + Y+G + + K +GL
 Sbjct: 12 VIFDGLTLDVDSSEPHYEAGRRTIARYGVDFPSWADHEAVVGISTQETVADWKRYYGLRA 71
 35 Query: 92 TVKEYIAENRRKRRQAVARDGVRPIKAGRLTHLHQHGYRLAVASSSPMVDIKRNLKEL 151
 TV+E +A NR ++AR R ++ + L G +AVAS S I L
 Sbjct: 72 TVEELLAVENRHYLGL-ARTSARAYPFEMRKVFELLAEGGVMAVAGSSPEIAAILART 130
 40 Query: 152 GVTECFEYMYVTGSDVSSSKPAFDPVFLRAAELLVDVDPKVCIVIEDTRHNGSLAAKAGMYC 210
 G+ +V+ +V+ KPARDVFL AA L +P C+V+ED G+ AA AARGC
 Sbjct: 131 GLDAHLRTVVSADAVARGKPAFDPVFLRAARLGTETPARCVVLEDPAAGAAAHAGMRC 189

- 45 A related DNA sequence was identified in *S. pyogenes* <SEQ ID 447> which encodes the amino acid sequence <SEQ ID 448>. Analysis of this protein sequence reveals the following:

Possible site: 25
 >>> Seems to have no N-terminal signal sequence

- 50 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3706 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

- 55 Identities = 62/202 (30%), Positives = 100/202 (48%), Gaps = 1/202 (0%)
 Query: 29 MEKVIIIFMDGVIVDSSEYTFLNKTEMLREBGI-DTDSYQYQYKITTPEFMQAMKEEFG 88

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M K I I F M D G V + D + E + L + + + + G I D + + G + + W + + +
 Sbjct: 3 MIKGIIFMDGVLFDTEPFYLRREDFPKTKGIPIDHLNSKDFIGNLQELMKELGNR 62
 Query: 89 LPKTVKEYIEMNNRRQAIVARDGVRPIKGAQRILHMLHQHYRLAVASSPMVDIKRNL 148
 5 VK + + +CA I + L + G + LAVAS + S D + L
 Sbjct: 63 DDAIVKAITTDYDAYEQAHKPPYQKLLITEVNSCLEQLKQGIKLAVASNSKRDQVLLAL 122
 Query: 149 KELGVTECFEYVMVTGEDVSSSKPAPDVPVRAARELLDVPKVCIVIEDTRNGSLAAKAGM 208
 + + + FE ++ EDVS KP PD + + A + L + K + V + ED + + G AAKAA +
 10 Sbjct: 123 ETTQIKDYFRIILAREDSRGKPYFDIYNKAVQKLSLQKQLLVVEDSQKGIATAAKANL 182
 Query: 209 YCFGFANPDYPPQOLSMADKVI 230
 F + Y D S AD I
 15 Sbjct: 183 TVFAITDYRY-GIDQSCADHKI 203

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 132

A DNA sequence (GBSx0138) was identified in *S.agalactiae* <SEQ ID 449> which encodes the amino acid
 20 sequence <SEQ ID 450>. Analysis of this protein sequence reveals the following:

Possible site: 20
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.22 Transmembrane 16 - 32 (16 - 32)
 25 ----- Final Results -----
 bacterial membrane --- Certainty=0.1086(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

30 The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 133

35 A DNA sequence (GBSx0139) was identified in *S.agalactiae* <SEQ ID 451> which encodes the amino acid
 sequence <SEQ ID 452>. Analysis of this protein sequence reveals the following:

Possible site: 34
 >>> Seems to have an uncleavable N-term signal seq
 40 INTEGRAL Likelihood = -5.04 Transmembrane 28 - 44 (27 - 45)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.3017(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 45 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 134

A DNA sequence (GBS01040) was identified in *S.agalactiae* <SEQ ID 453> which encodes the amino acid sequence <SEQ ID 454>. Analysis of this protein sequence reveals the following:

```

Possible site: 17
5  >>> Seems to have an uncleavable N-term signal seq
    INTEGRAL    Likelihood =-10.72    Transmembrane    38 - 54 ( 34 - 60)
    INTEGRAL    Likelihood = -7.70    Transmembrane    4 - 20 ( 1 - 22)
    INTEGRAL    Likelihood = -4.99    Transmembrane    153 - 169 ( 150 - 171)
10  INTEGRAL    Likelihood = -2.55    Transmembrane    179 - 195 ( 178 - 198)
    INTEGRAL    Likelihood = -2.39    Transmembrane    93 - 109 ( 93 - 109)
    INTEGRAL    Likelihood = -1.17    Transmembrane    116 - 132 ( 116 - 133)
    INTEGRAL    Likelihood = -0.43    Transmembrane    344 - 360 ( 344 - 360)

----- Final Results -----
15  >>> bacterial membrane --- Certainty=0.5288 (Affirmative) < succ>
    bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
    bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

20  >GP:CAB14853 GB:Z99118 two-component sensor histidine kinase
    [Bacillus subtilis]
    Identities = 254/585 (43%), Positives = 371/585 (63%), Gaps = 9/585 (1%)

Query: 2  LMVLLFQRLGIIMILAFLLVNNYSFROLIEERSK-RETVVLVIIPGLFVLIISNITGIEIK 60
15  1M+++ +R+GII+IL F+L + FRQ ++ + + +L+ IF LF IISN TGIEI+
    Sbjct: 4  LMIMGLERVGIIIVILGFIHAHTKLPQALQNDQGYGKAILISIFSLPSISINVTGSIQ 63

Query: 61  GDRSLVERPFLTITISHSOSLANTKTIVITIASLVGGPLVGSIVGPIGGVHRFFQGSFSGS 120
30  + +V ++ TI S S+ANTR L + L+GGP VG+ +G + G+HRF G +
    Sbjct: 64  RNM-IVNNVWVFITDPGSGTANTRILGIVEIGGLGGPFVGAGIGLALGHRFSLGGSTL 122

Query: 121 FYIVSSVLVGIVSGKIGDKLKENHLYPSTQVILLISIIAESIQMLFVGIFT-----GWEL 175
    VSS+L G+++G IG + + P+ L+ I R+Q+ + + WEL
    Sbjct: 123 SCAYSSILAGVLAGLIGRYFTKRYMPTPRIAALVGIGMESLQMIILLMAKPPSDAWEL 182

35  Query: 176 VKMIVIPMILINSLSGSLFLAILKTYLSNESQLRAVQTRDVLSEITROTLFYLROGLTPQS 235
    V MI IPM+++N GS +FL+I++ + E Q RA++T VL + QTLP+ RQGL S
    Sbjct: 183 VSMIGIFMILINTGSGFIFLSIIQAIIRKEQARALETHRVLTIADOTLFFPFRQLNENS 242

40  Query: 236 ARSVCEIIRKTNFNDVAVGLTDRSNVLAHIGVGDHHAAGQPVKTDLSKSVIFDGEPIRAQ 295
    +SV II + T DAV LTD+ +LH+G G DHHI + + T LSK VI G A
    Sbjct: 243 CKSVAALIHKLGTDAVSLTDKELIAHVAGMDHHPKSKLITGLSKKVIKTHIMIAL 302

Query: 296 DKAISCPDHNCQNSAIVVPLKINDKTGALKMYFAGDKTMSVEVENILVLGLAQIFSGQ 355
45  + I C C L++ATV+PL N T+G LKMYF +S+VER L GLA +PS Q
    Sbjct: 303 SQEEIECTHACPLHAATVILPLTNSNGTITGLKMYFKPSAGLSQVREELAGLMLFSTQ 362

Query: 356 LAMGITEBQNKLAGMAEIKALQAQINPHFFNAINTISALIRIDSDIKARYALMQLSTFFR 415
    L +G E Q+KL ARKALQAQ+NPHF PNAINTISAL R D +K R L+QLS +FR
50  Sbjct: 363 LEIGEAELQKLLGDARAKALQAQVNHPLFNAININTISALCRTDVEKTRKLLQLSVFTR 422

Query: 416 TSLQGGQREVTLRQEKSHVDAYMNVKELFPDKYQLSYDI -SAPENKLEPPFGLQLVLE 474
    ++LQG + + L +S +H+AY++R+ RFP KY++ +I S E++++PPF LQLVLE
55  Sbjct: 423 SNLQGARQQLLPLSKRLNHLNAYLSLQARFPFGKYKILNIDSRIGTETIPFVLQLVLE 482

Query: 475 NAVRHAPKERTDNHILLVOIKPDORHYCVSVSDNGQGISDTIIDLQOETVAESKGTGTA 534
    NA+REAF +++ + V + D + V+DNG+GI ++ +IG++ +GTGTA
    Sbjct: 483 NALREAFPKKQDICKVTVCVLSDDASVMKVDNRGCIIPDVLPLGAKKPPFSKEGTGTA 542

60  Query: 535 LVNLNRLNLLYGSVSLHFSSD -KNGTKVWYRIPNRIREDEHN 578
    L NIN RL L+G + LH SS+ GT+V +++P + ++ R+
    Sbjct: 543 LVNLNRLTLGLPQQALHISSEVHRGTETVSPQVPMQMKRGEH 587

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 455> which encodes the amino acid sequence <SEQ ID 456>. Analysis of this protein sequence reveals the following:

```

Possible site: 23
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.1771(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

```

Identities = 75/245 (30%), Positives = 117/245 (47%), Gaps = 22/245 (8%)

Query: 348 LAQIFSOQL----AMGITEEONKLASMAEIKALQAQINPHFFFNAINAISALIRI-DSD 401
      LAQ F+ L M ++K ++AL+QINPEF +N ++TI + DS
Sbjct: 4 LAQQFNALLDQIDGLMVAADKEKALGQYELQALASQINPHFLYNTLDTIIMAEFNDK 63

Query: 402 KARYALMQLSTFFRTSLQGGQDREVILQEOKSHVDAYMNVKLRFPDKYQLSYDISAPE- 460
      + L+ +PR +L G + + L E HV Y+ ++K R+ DK LSY++ +
Sbjct: 64 RVVEVTKSLAKYFRALNAGNEY-IRLADELHVSQYLFIQKQRYGDK--LSYEVQGLDV 120

Query: 461 --KMKLPPFGLQVLVENA+RHAFKERKTDNHILWQIKPDGHYYCVSVSDNGQGISDTIID 518
      +P LQ LVENA+ H KE I V + + ++V DNG+GI D+ +
Sbjct: 121 YADFVPIKLLQLVLENATYHGIKEVDKRGMIKVTVDIAQHLMLTVMDNGKGIETDSSLT 180

Query: 519 KLGQETVAESKGTGTALVNLNRLNLLYGS--VSLHFSSDKNGTKWYVPRN---IRE 573
      Q +A G L N++ RL L YG +H SD+ T++ +P + +
Sbjct: 181 N-SQSLIARG---GVGLKNVDQRLKLHYGEGYHMTIHSQSDQ-FTETQLSLPKMHLEMD 235

Query: 574 DEHEN 578
      D EN
Sbjct: 236 DTQEN 240

```

SEQ ID 454 (GBS248d) was expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 124 (lane 2-4; MW 71kDa). It was also expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 124 (lane 5-7; MW 46kDa) and in Figure 180 (lane 2; MW 46kDa).

GBS248d-His was purified as shown in Figure 234, lane 3-4.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 135

A DNA sequence (GBSx0141) was identified in *S.agalactiae* <SEQ ID 457> which encodes the amino acid sequence <SEQ ID 458>. This protein is predicted to be two-component response regulator (lytT). Analysis of this protein sequence reveals the following:

```

Possible site: 61
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3230(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 9495> which encodes amino acid sequence <SEQ ID 9496> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

- >GP:CAB14852 GB:Z99118 two-component response regulator [Bacillus subtilis]
Identities = 105/244 (43%), Positives = 157/244 (64%), Gaps = 6/244 (2%)
- 5 Query: 3 MKILLDEMFARQELSLVHVSQVDMPEIQAEDISEAKKILFRQOIDLIFDISLSE 62
+++LI+DEEM AR EL++L++ + D EI +AE+I A + Q+ DL+FLDA LS
Sbjct: 2 LRVLVDDERLARDELAYLLKTN--DEMEINEAENIESAFQNMDDQKPLLFLVDLSS 59
- 10 Query: 63 ENGFTLANQLSQLAHPPLAVFATYDNYAVKAFESNAVVDYIMKPFQQRVDMLSKVK 122
ENG F +A +L ++ H P +VFATYD Y A+K A F +A+D Y+ K P F++R+ L K R K+
Sbjct: 60 ENGFDIAKRLKMKHPPIAVFATYDQYALKAFFEDALDYLT K P FDEIRIQCTLGKYK 119
- 15 Query: 123 SQLTTASDVQAIPKASVELLTITLSRSVVVMQDIVAASVEDGELTVSTVQKTYTR 182
++ VE A L L++ + V+V +DI+ A EDG + V T +YT+
Sbjct: 120 NR----DIVTEQNSHAGQKIALSVGESIVIVTKDIIYAGTEDGHVNVKPFDSHYTS 175
- 20 Query: 183 KTLNWPFSRAVAPYPLQIHRTIVNLEMIIEIQPFNHTLLMSNGKFPVGRSLKDL 242
TL ++ + F++HR+ V+V E I+EIQPFN T LIM +G K PV +Y K+L
Sbjct: 176 DTLAVVIEKKLPDSDFIRVRSFVVNTEYIKKIQPFNFTYNLIMDKSGKIPVSTYAKEL 235
- 25 Query: 243 NEHL 246
+ L
Sbjct: 236 KKLL 239
- 25 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 459> which encodes the amino acid sequence <SEQ ID 460>. Analysis of this protein sequence reveals the following:
- Possible site: 27
>>> Seems to have no N-terminal signal sequence
- 30 ----- Final Results -----
bacterial cytoplasm --- Certainty=0.3818 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
- 35 An alignment of the GAS and GBS proteins is shown below:
- Identities = 44/148 (29%), Positives = 84/148 (56%), Gaps = 5/148 (3%)
- Query: 5 LILLDEMFARQELSLVHVSQ-EVDMPEIQAEDISEAKKILFRQOIDLIFDISLSE 63
+LI++DE RQ + LV+ SQ +D + +AE+ A + ++ D++ DI++ +
Sbjct: 4 LLIVEDEYLVRQIRSLVDPSQPKIDR--VNEAENGQLANDLFQKPEYDVIITDINPKL 61
- 40 Query: 64 NGFTLANQLSQLAHPPLAVFATYD--NYAVKAFESNAVVDYIMKPFQQRVDMLSKVK 121
NG LA + Q + +VF T YD NYA+ A + A DY++K P F + V+ L K +K
Sbjct: 62 NGIQALIELKQSPQTHLVLTGYDDFNIALSALKLGADDYLLKPFSAVDENMLKLRK 121
- 45 Query: 122 LSQLTASDVQAIPKASVELLTITLS 149
+L+ ++ Q + ++ E+ + ++
Sbjct: 122 KLELSKRTTETIQELVQPKVEALAMA 149
- 50 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 136

A DNA sequence (GBSx0142) was identified in *S.galactiae* <SEQ ID 461> which encodes the amino acid sequence <SEQ ID 462>. Analysis of this protein sequence reveals the following:

- 55 Possible site: 18
>>> Seems to have no N-terminal signal sequence
- Final Results -----
bacterial cytoplasm --- Certainty=0.0266 (Affirmative) < succ>

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```

bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

- 5 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 137

A DNA sequence (GBSx0143) was identified in *S.agalactiae* <SEQ ID 463> which encodes the amino acid sequence <SEQ ID 464>. Analysis of this protein sequence reveals the following:

```

Possible site: 37
>>> Seems to have no N-terminal signal sequence
INTEGRAL    Likelihood =-11.89    Transmembrane    104 - 120 ( 99 - 134)
INTEGRAL    Likelihood = -5.89    Transmembrane    47 - 63 ( 46 - 65)
15 INTEGRAL    Likelihood = -3.29    Transmembrane    22 - 38 ( 21 - 39)
INTEGRAL    Likelihood = -2.81    Transmembrane    74 - 90 ( 70 - 92)

----- Final Results -----
20 bacterial membrane --- Certainty=0.5755 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 8499> which encodes amino acid sequence <SEQ ID 8500> was also identified.

- 25 The protein has homology with the following sequences in the GENPEPT database:

```

>GP:CAB14851 GB:Z99118 similar to hypothetical proteins from B. subtilis [Bacillus
subtilis]
Identities = 50/110 (45%), Positives = 82/110 (74%), Gaps = 2/110 (1%)

30 Query: 20 QMSIYAAILLVSNQIMSMLEKSLPIPTTVIGLVIMYVLLTAKIKVWVDSFGALMISM I 79
Q I+I I+IWS MI+ ++P +PIP +V+GLVL+++LL K+IK+E V++ G + S+I
Sbjct: 12 QAPIFAVIMVLSNMIAIVP--IPIPASVVGVLVLLFLLCLLKVKIKLEQVETIGTSLTSLI 69

Query: 80 GFMFVPSGISVAANLQILKAEGLQVLAVITISTVMMVLVVVYVARLILAI 129
GF+FPVPSGISV +L +++ GLQ+V VI ++T+++L ++LIL++
35 Sbjct: 70 GPLFVPSGISVMNSLGMQCYGLQIVLVILLATITILLGATGLFSLQLSL 119

```

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 138

A DNA sequence (GBSx0144) was identified in *S.agalactiae* <SEQ ID 465> which encodes the amino acid sequence <SEQ ID 466>. Analysis of this protein sequence reveals the following:

```

Possible site: 44
>>> Seems to have a cleavable N-term signal seq.
INTEGRAL    Likelihood =-12.21    Transmembrane    219 - 235 ( 208 - 241)
INTEGRAL    Likelihood =-11.94    Transmembrane    103 - 119 ( 99 - 133)
INTEGRAL    Likelihood = -5.57    Transmembrane    157 - 173 ( 154 - 175)
50 INTEGRAL    Likelihood = -1.70    Transmembrane    73 - 89 ( 73 - 89)

----- Final Results -----

```

bacterial membrane --- Certainty=0.5883 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

5 The protein has homology with the following sequences in the GENPEPT database:

>GP:CAB14850 GB:Z99118 similar to hypothetical proteins [Bacillus subtilis]
 Identities = 120/240 (50%), Positives = 159/240 (66%), Gaps = 10/240 (4%)

10 Query: 1 MELLATPIPGICFSLILYITIGSEHLFKKSGKFFLLQPLFFAMVSGIVILWMSKGLGTDVK 60
 ME +F FGI SL + IG LFKK+KGFLL PLF AMV GI L +
 Sbjet: 1 MBSIMSIPFGIVISLAAPFGIGTFLFKKIKGFFLLPLFVAMVLGLAFL- - - - - KIG 51

15 Query: 61 TPTQAYKPGGDLIFWFLNPATIAFAVFLYKKNDVVKVWELLSSLVIGMIVSLILIVA 120
 F Y GG++I +FL PATIAFA+FLYK+ D +KKYF +I++S+ G I S+ ++
 Sbjet: 52 GFSYADIRNGGELIKFFLEPATIAFAIPLFYKQDKLKKYVQIMASIIAGSICSVTIVYL 111

20 Query: 121 ISKVKGLSCVGIASMLPQAATTALPATAIGGNTAVTMACILNAVIIYALGKKLVSF 180
 ++K + L + SFLPQAATTALP++ IGG + TA A I NAVI+YALG
 Sbjet: 112 LARGHLDSAVMKSMPLPQAATTALALFLKGIIGISDITAFAVIFNAVIVYALGALFLK 171

25 Query: 181 FHLNDSKIGAGLGLTSGHTVGAAPALELGEIQGMAIAVTVVIGLVVDLVPIPSHLIG 240
 F + + I GL LSTSGH +G A +B+GE++ AWA+IAVVV+G+V LVIP+F LIG
 Sbjet: 172 FKVK-NPISKGLALGTSCHALGVAVGIEGVEANASIAVVVGVVTVLVIPVVFQLIG 230

25 No corresponding DNA sequence was identified in *S. pyogenes*.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 139

A DNA sequence (GBSx0145) was identified in *S. agalactiae* <SEQ ID 467> which encodes the amino acid sequence <SEQ ID 468>. Analysis of this protein sequence reveals the following:

Possible site: 22
 >>> May be a lipoprotein

----- Final Results -----

35 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

40 Identities = 508/542 (93%), Positives = 523/542 (95%)

Query: 1 MTKYLKYSFVALFLASIFLVACQNNQSTKIKRTRKQRPKDELVVSNGAKLPHFDPKDR 60
 ++KYLK Y S + LFL + LVACQ Q CTKER RKQRPKDELVVSNGAKLPHFDPKDR
 45 Sbjet: 3 VSKYLKYFSITLFLTGILVACQQKQCTKERQRKQRPKDELVVSNGAKLPHFDPKDR 62

Query: 61 YGHNENITHSTLLKRSPELDIKGLAKYKISKDGLTWSFDLHDDFKPSNHEPVTADD 120
 YG+HNENITHSTLLKRSPELDIKGLAK Y +S+DGLTWSFDL+DDFKPSNHEPVTADD
 50 Sbjet: 63 YGVHNEGITHSTLLKRSPELDIKGLAKYKISDGLTWSFDLHDDFKPSNHEPVTADD 122

Query: 121 VKPTYMLKADGKAWDLTFIKNVSVVGKNQVNIHLTEAHSTFTAGLTIPIVVKHYNDK 180
 VKPTYMLKADGKAWDLTFIKNVSVVGKNQVNIHLTEAHSTFTAGLTIPIVVKHYNDK
 55 Sbjet: 123 VKPTYMLKADGKAWDLTFIKNVSVVGKNQVNIHLTEAHSTFTAGLTIPIVVKHYNDK 182

Query: 181 YKSNFIGSGPYMVVKRYKAGBQAI FVRNPFYHKKPKYFKGWTWLLDENTALAALSGDVD 240
 YKSNFIGSGPYMVVKRYKAGBQAI FVRNPFYHKKPKYFKGWTWLLDENTALAALSGDVD
 60 Sbjet: 183 YKSNFIGSGPYMVVKRYKAGBQAI FVRNPFYHKKPKYFKGWTWLLDENTALAALSGDVD 242

Query: 241 MIYATPELASKVKYKTRLLDIASNDVRLSLPYVKKGVKNSPDQYVGVNDVSDPAIRK 300
 MIYATPELA KVKYKTRLLDI SNDVRLSLPYVKKGV+ +SPDQYVGVNDVSDPAIRK

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Sbjct: 243 MIYATPELADKKVGRLLDIPSRDVRGLSLPYVKKGVITDSDPGYFVGNVDYEDPAIRK 302

Query: 301 ALITGLNRQKVLDTVLNLYGKPAYSTIDRTPFNNKTAIDKNKVAKQLLTKAGWKEQA 360
 5 Sbjct: 303 ALTIGLRQKVLDTVLNLYGKPAYSTIDKTFFNNKTAIDKNKVAKQLLTKAGWKEQA 362

Query: 361 DGSRRKKNLKSDFLYPTNDQLRANLAVEVAQKALGTTIKLQASNDWMATKSHDSA 420
 10 Sbjct: 363 DGSRRKKNLKSDFLYPTNDQLRANLAVEVAQKALGTTIKLQASNDWMATKSHDSA 422

Query: 421 LLYAGGRHHQQFYESHYPBLAGKSWNITFPYNNFTVTKYLDKAMTSPDLKAMKWKLA 480
 15 Sbjct: 423 LLYAGGRHHQQFYESHYPBLAGKSWNITFPYNNFTVTKYLDKAMTSPDLKAMKWKLA 482

Query: 481 QWDGKTCASTLGLDFNFWLVLNHTTYIGDKRINVCKQGVISHGHWSLLTNIAEWTWDES 540
 20 Sbjct: 483 QWDGKTCASTLGLDFNFWLVLNHTTYIGDKRINVCKQGVISHGHWSLLTNIAEWTWDES 542

Query: 541 AK 542
 25 Sbjct: 543 TK 544

There is also homology to SEQ ID 60.

A related GBS gene <SEQ ID 8501> and protein <SEQ ID 8502> were also identified. Analysis of this
 25 protein sequence reveals the following:

Lipop: Possible site: 22 Crend: 5
 MoG: Discrim Score: 10.46
 GVH: Signal Score (-7.5): -1.29
 Possible site: 22
 30 >>> May be a lipoprotein
 ALOM program count: 0 value: 7.27 threshold: 0.0
 PERIPHERAL Likelihood = 7.27 386
 modified ALOM score: -1.95

35 *** Reasoning Step: 3

----- Final Results -----
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 40 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

SEQ ID 8502 (GBS106) was expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell
 extract is shown in Figure 18 (lane 3; MW 61kDa).

The GBS106-His fusion product was purified (Figure 19A, lane 2) and used to immunise mice. The
 45 resulting antiserum was used for Western blot (Figure 255A), FACS (Figure 255B), and in the *in vivo*
 passive protection assay (Table III). These tests confirm that the protein is immunoaccessible on GBS
 bacteria and that it is an effective protective immunogen.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for
 vaccines or diagnostics.

50 Example 140

A DNA sequence (GBSx0146) was identified in *S. galgaliae* <SEQ ID 469> which encodes the amino acid
 sequence <SEQ ID 470>. Analysis of this protein sequence reveals the following:

Possible site: 41
 >>> Seems to have no N-terminal signal sequence

55

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----- Final Results -----

bacterial cytoplasm --- Certainty=0.4862(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

5

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

10 **Example 141**

A DNA sequence (GBSx0147) was identified in *S.galactiae* <SEQ ID 471> which encodes the amino acid sequence <SEQ ID 472>. Analysis of this protein sequence reveals the following:

Possible site: 19

>>> Seems to have no N-terminal signal sequence

15

INTEGRAL	Likelihood = -7.27	Transmembrane	252 - 268 (249 - 275)
INTEGRAL	Likelihood = -5.73	Transmembrane	67 - 83 (62 - 90)
INTEGRAL	Likelihood = -5.26	Transmembrane	107 - 123 (104 - 134)
INTEGRAL	Likelihood = -3.77	Transmembrane	153 - 169 (152 - 170)

20

----- Final Results -----

bacterial membrane --- Certainty=0.3909(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

25

A related GBS nucleic acid sequence <SEQ ID 9295> which encodes amino acid sequence <SEQ ID 9296> was also identified.

The protein differs from U78968 at the N-terminus:

Query: 1 MASVNYDTSLTPVQYKAIAHHYGLDKPAPVQYFIWLKNIQGHGSLVYRQPFVIDIIRS 60
 MASVNYDTSLTP QYKAIAHHYGLDKPA VQYFIWLKN IQG IGTSLVYRQPV DIIRS
 30 Sbjct: 39 MASVNYDTSLTPAQYKAIAHHYGLDKPALVQYFIWLKNIQGHGSLVYRQPFVSDIIRS 98

30

There is also homology to SEQ ID 64.

A related GBS gene <SEQ ID 8471> and protein <SEQ ID 8472> were also identified. Analysis of this protein sequence reveals the following:

35

Lipop: Possible site: -1 Crend: 10
 MoG: Discrim Score: 3.72
 GVH: Signal Score (-7.5): -5.37
 Possible site: 40

40

>>> Seems to have an uncleavable N-term signal seq

ALOM program	count: 5	value: -7.27	threshold: 0.0
INTEGRAL	Likelihood = -7.27	Transmembrane	290 - 306 (287 - 313)
INTEGRAL	Likelihood = -5.69	Transmembrane	12 - 28 (11 - 33)
INTEGRAL	Likelihood = -5.73	Transmembrane	105 - 121 (100 - 128)
INTEGRAL	Likelihood = -5.26	Transmembrane	145 - 161 (142 - 172)
INTEGRAL	Likelihood = -3.77	Transmembrane	191 - 207 (190 - 208)
PERIPHERAL	Likelihood = 2.97		245
modified ALOM score: 1.95			

45

*** Reasoning Step: 3

50

----- Final Results -----

bacterial membrane --- Certainty=0.3909(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

SEQ ID 8472 (GBS436) was expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 173 (lane 9; MW 54kDa).

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 142

A DNA sequence (GBSx0148) was identified in *S.galactiae* <SEQ ID 473> which encodes the amino acid sequence <SEQ ID 474>. This protein is predicted to be transmembrane transport protein DppC (oppC). Analysis of this protein sequence reveals the following:

```

10 Possible site: 39
    >>> Seems to have a cleavable N-term signal seq.
        INTEGRAL Likelihood = -8.28 Transmembrane 77 - 93 ( 68 - 101)
        INTEGRAL Likelihood = -7.80 Transmembrane 182 - 198 ( 180 - 204)
15 INTEGRAL Likelihood = -7.06 Transmembrane 112 - 128 ( 104 - 132)
    INTEGRAL Likelihood = -5.10 Transmembrane 239 - 255 ( 235 - 258)

    ----- Final Results -----
        bacterial membrane --- Certainty=0.4312(Affirmative) < succ>
        bacterial outside --- Certainty=0.0000(Not Clear) < succ>
20 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

There is homology to SEQ ID 68.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 143

25 A DNA sequence (GBSx0149) was identified in *S.galactiae* <SEQ ID 475> which encodes the amino acid sequence <SEQ ID 476>. This protein is predicted to be ATPase protein DppD. Analysis of this protein sequence reveals the following:

```

30 Possible site: 59
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
        bacterial cytoplasm --- Certainty=0.1957(Affirmative) < succ>
        bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
35 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein differs from U78968 at the C-terminus:

```

Query: 241 QTSEFARSLWRSLAPQQRFLKGVTHDLRG 267
      QTSEFAR LWR+LFQQ+FLKGVTHDLRG
30 Subjct: 241 QTSEFARRLWRSLAPQQRFLKGVTHDLRG 267

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 477> which encodes the amino acid sequence <SEQ ID 478>. Analysis of this protein sequence reveals the following:

```

45 Possible site: 59
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
        bacterial cytoplasm --- Certainty=0.1957(Affirmative) < succ>
        bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
50 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

Identities = 255/267 (95%), Positives = 262/267 (97%)

```

5  Query: 1  NTEVTLSSIKDLSITFTQYGRFLKPPQSTPIQALNLKIKKGLLAIGASGSGKSLLAHAI 60
   Sbjct: 1  NTEVTLSSIKDLSITFTQYGRFLKPPQSTPIQALNLKIKKGLLAIGASGSGKSLLAHAI 60

10 Query: 61  NDILPKNAASVIGDMITYRGQSLMSKRIRKQLRGKQITLIPQSVNYLDPSITVKHKVRLGISSE 120
   Sbjct: 61  NDILPKNAASVIGDMITYRGQSLMSKRIRKQLRGKQITLIPQSVNYLDPSITVKHKVRLGISSE 120

15 Query: 121  NSKATQEGFLPQQFGKLGESDGLYFPQLSGGMLRRVLPFTTCISDKVSLIIADEPTPLGLHFD 180
   Sbjct: 121  NAKATQEGFLPQQFGKLGESDGLYFPQLSGGMLRRVLPFTTCISDVSLIIADEPTPLGLHFD 180

20 Query: 181  ALQMVLQQLRSFADKGISVIFITHTDIVAASQIADRIITIPKEKKAETAPASFFSGGGEQL 240
   Sbjct: 181  ALQMVLQQLRSFADKGISVIFITHTDIVAASQIADRIITIPKEKKAETAPASFFSGGGEQL 240

25 Query: 241  QTEFARSLWRSLPQQFGFLKGVTHDLRG 267
   Sbjct: 241  QTEFARSLWRSLPQQFGFLKGVTHDLRG 267

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 144

A DNA sequence (GBSx0150) was identified in *S. agalactiae* <SEQ ID 479> which encodes the amino acid sequence <SEQ ID 480>. This protein is predicted to be ATPase protein DppE. Analysis of this protein sequence reveals the following:

```

30 Possible site: 41
   >>> Seems to have no N-terminal signal sequence

   ----- Final Results -----
35   bacterial cytoplasm --- Certainty=0.3783 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 481> which encodes the amino acid sequence <SEQ ID 482>. Analysis of this protein sequence reveals the following:

```

40 Possible site: 41
   >>> Seems to have no N-terminal signal sequence

   ----- Final Results -----
45   bacterial cytoplasm --- Certainty=0.3383 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

Identities = 188/205 (91%), Positives = 197/205 (95%)

```

50 Query: 1  NTLSEAKKLGIFYHKDKQWLFEKINLEVAAPQVIGIFGQSGCGKTSLSRVLAGFLHPKSGEV 60
   Sbjct: 1  NTLSEAKKLGIFYHKDKQWLFEKIDLEVAAPQVIGIFGQSGCGKTSLSRVLAGFLHPKSGEV 60

55 Query: 61  LVDGSHLPSKAFRPVQLIQHPEKTMNLEWPKKSLSEAYYPSRLIDAFGIQEKWLRR 120
   Sbjct: 61  LVDGSHLPSKAFRPVQLIQHPEKTMNLEWPKKSLSEAYYPSRLIDAFGIQEKWLRR 120

```

Query: 121 PSELGGGLQRFPSVRSIHPTKYLIADEMTIMLDGITOASVWKSLLSEIVKDRNLGLIVI 180
 PSELGGGLQRFPSIVRSIHPTKYLIADEMTIMLDGITOASVWKSLLSEIVKDRNLGLI+I
 Sbjct: 121 PSELGGGLQRFPSIVRSIHPTKYLIADEMTIMLDGITOASVWKSLLSEIVKDRNLGLIII 180

5 Query: 181 SHDFMLEKLCNQCYMIEENRIVSF 205
 SH+F MLEKLC+ CYMIEENR F
 Sbjct: 181 SHEFMLEKLCNQCYMIEENRIVOLF 205

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 145

A DNA sequence (GBSx0151) was identified in *S. agalactiae* <SEQ ID 483> which encodes the amino acid sequence <SEQ ID 484>. This protein is predicted to be PTS system, trehalose-specific IIBC component (treB). Analysis of this protein sequence reveals the following:

Possible site: 59
 >>> Seems to have no N-terminal signal sequence

INTEGRAL	Likelihood = -10.14	Transmembrane	468 - 484 (462 - 489)
INTEGRAL	Likelihood = -8.23	Transmembrane	279 - 295 (275 - 306)
INTEGRAL	Likelihood = -6.05	Transmembrane	112 - 128 (105 - 130)
INTEGRAL	Likelihood = -3.35	Transmembrane	204 - 220 (203 - 222)
INTEGRAL	Likelihood = -1.75	Transmembrane	255 - 271 (255 - 271)
INTEGRAL	Likelihood = -1.54	Transmembrane	327 - 343 (326 - 344)
INTEGRAL	Likelihood = -0.37	Transmembrane	422 - 438 (422 - 438)
INTEGRAL	Likelihood = -0.06	Transmembrane	304 - 320 (304 - 320)

----- Final Results -----

bacterial membrane	---	Certainty=0.5055 (Affirmative)	< succ>
bacterial outside	---	Certainty=0.0000 (Not Clear)	< succ>
bacterial cytoplasm	---	Certainty=0.0000 (Not Clear)	< succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAF94072 GB:AB004175 PTS system, trehalose-specific IIBC
 component [Vibrio cholerae]
 Identities = 225/484 (46%), Positives = 318/484 (65%), Gaps = 28/484 (5%)

Query: 5 KHDKALLLEAIGKENISAVTHCATRMRFVLNDSSKAKVKVIELPSVKGITFTNAQQPQV 64
 K D L+E +GG+ NI++VTEC TR+RPVLN +A +E L VKG FTNAQQPQV
 Sbjct: 10 KQDVTRILVVGESNIASVTHCLTRFLVNLQPEQADKAGLEALSMVKGCFPTNAQQPQV 69

Query: 65 IIGNDVPIFYNAFVAVSGIEGVSKENAKSAQKQNFQLRVLMIAEFTPIIPAIIVGG 124
 +IG +V Y + +G +VSK+ AK AA++N N L+R ++ LRIIF P++PAII VG
 Sbjct: 70 VIGTEVDQVYKMLEQTGKQAVSKDKDAKVAARQNMNVLERGISHLARIFVPLLPATITGG 129

Query: 125 LILGFERNILDVAFPEFLQCKVVDGVRQVDSGGSHPIWNTLVDVSTFMSGVDSFLMLPGEAI 184
 LILGFERN++ ++ DG TL ++S FW+ V +FLML GEAI
 Sbjct: 130 LILGFERNVIGDI-----RMFDG-----KTLTETISQWASVHAFNLIGRAI 170

Query: 185 FHFPLVGIIVSVIRKMGITQILGIVLIGLCLVSPQLLNAYSVASTSAADIAKQWNSWIFGYF 244
 F FLVFG+ WS +K+G T ILGL LG+ LVSEQL+NAY + W+FG F
 Sbjct: 171 FFFFLVGVGVSTVKKLGCTFLIGLITGLVTLVSPQLMNAYLIGKEVPE-----VWDFGLF 224

Query: 245 TVQKIGYQAVQVIFALLAGLSLSYLRIEPRKHIPEVVSIMFVPLSLVPALIAHTVLOPI 304
 ++K+GYQAVQVIPA+LAG+L++E R+ +P ++ VPP+S++ ++LH+ +GP
 Sbjct: 225 AIEKVGQYQAVQVIFALLAGVALAFENNLRKRVPSYLVVVVPPVSITVSVLNAFPIGPF 284

Query: 305 GWTLGKVISAIVLIGLTPGVKMLPGAIFGALYAPPVITGLHMTNAIDTQLIADTRKTH 364
 G +G ++ +TG ++ +FG +YAP VITG+HH TNA+D QL+ + T
 Sbjct: 285 GRVIGDGVAFPAKNAITGDFAVIGSTLFGEMYAPLVITGIHHTTNAVDLQMQE--LGST 342

Query: 365 GLWPMIALSNIAQGSVALVYFMHREHDEKSAQISLPAISAYLVGTEPAFGVNVKVIYP 424
 +WP+IALSNIAQ SAV+ ++ + E IS+PAISAYLVGTEPA++G+N+KY +P

-220-

Sbjct: 343 PIWPLIALSNIAQASAVVGIIIIISK-KQGERDISVPAAISAYLVGTEPMYGINLKYKFP 401

Query: 425 FVAGMIGSSVAGLLATTENNQANSIGVGGLPGFTLINVKYMGFFIOANVAIFPLPLTY 484
++ MIGS+AA + + V AN IGVGSLG LSI ++ + + M +AI +P LTL

Sbjct: 402 MLSAMIGSALAAAVCSAGVMANGIGVGGLPGILSIQPFWSIYLVNMLIAILVPAALTY 461

Query: 485 FFKK 488

K

Sbjct: 462 LMYK 465

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 485> which encodes the amino acid sequence <SEQ ID 486>. Analysis of this protein sequence reveals the following:

Possible site: 59

>>> Seems to have no N-terminal signal sequence

INTEGRAL Likelihood = -9.61 Transmembrane 466 - 482 (457 - 488)

INTEGRAL Likelihood = -8.01 Transmembrane 279 - 295 (275 - 306)

INTEGRAL Likelihood = -6.05 Transmembrane 112 - 128 (105 - 130)

INTEGRAL Likelihood = -3.35 Transmembrane 204 - 220 (203 - 222)

INTEGRAL Likelihood = -3.13 Transmembrane 255 - 271 (255 - 272)

INTEGRAL Likelihood = -2.07 Transmembrane 327 - 343 (325 - 344)

INTEGRAL Likelihood = -0.59 Transmembrane 422 - 438 (422 - 438)

----- Final Results -----

bacterial membrane --- Certainty=0.4843(Affirmative) < succ>

bacterial outside --- Certainty=0.0000(Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP:AAF94072 G3:AE004175 PTS system, trehalose-specific IIBC

component [Vibrio cholerae]

Identities = 231/484 (47%), Positives = 322/484 (65%), Gaps = 28/484 (5%)

Query: 5 EQDANSLTAIGSKENIKVTHCATMRFLVNDNNKANVKEIEKISVVEGTFPNAQQPV 64

+QD L+ +GG+ HI VTEC TR+RFVLN ++ +E +S+VSG FDNAGQPV

Sbjct: 10 KQVTRILTELVOGESNTASVTHCLTRLRFTVNAQPEQADKAGLSMSVGCFTNAGQPV 69

Query: 65 IIGNDVPFVNDFTAVSSIEGVSKAAKSAKSNQALQVMTLARIPTLIPALIVG 124

+IG +V Y + + VSK+ AK AA+ N N L+R ++ LAEIF P++PAII GQ

Sbjct: 70 VIGTEVDQVYIMLLEQTKKQAVSKDDAKVAARQNNVLERGISHLAEIFVPLLPALITGG 129

Query: 125 LILGFNNILSFVPEFLAQCVKSGKLNFDAGDPVNTIVKVSPPNSGVNHFMLAGEAT 184

LILGFNN++ + +FD T+ +S FW+ V+ FIML GRAT

Sbjct: 130 LILGFNNVIGDI-----RMFDG-----KYLTFISQFNASVHAFMLIGRAI 170

Query: 185 FHEFLVQITVSVTRMGITQLIGLVIGICLVSPQLINAVAVAGTFAARIKNNWDFQFP 244

F FLELVG+ WS +K+G T ILGI LG+ LVNSQL+HAY + G E WDFG F

Sbjct: 171 FFFFLVGVCMSTVKLGSTPIIGITLTVLSPQLMGAYLI-GKEVPE-----WDFGLF 224

Query: 245 TIDRIGYQAVIPALLAGLSLAYLPIFWKRIRPEVVSIMFVPHSLIPALITAMTGLPI 304

I ++GYQAVIP+LAG+LAI+E R+ P + + VEP+SAI ++LAIH +GP

Sbjct: 225 AIEKVGYQAVIPAILAGVALAFTENNLRVVPSYLYLVVFPVSITVSVLHAFTGIFP 284

Query: 305 GWITIGNISFVVLAGLTGPMVLMFGALPGALYAPLVITGLHMMNIAIDTOLIAETATKTT 364

G IG G+AF A +TG + +FG +YAPLVITG+HH TRAAD QL+ + T

Sbjct: 285 GRVIGDGAFAAKAMTSDFAVIGSTLPGFMVAPLVITGIHHTNAVLDLQMSLG--GT 342

Query: 365 QINPITALSNIAQGSAYFAYILNRRHEKREARISLPAAISAYLVGTEPALGVNFKVYVP 424

+NP+IALSNIAQ SAV ++++ ++ E +IS+PAISAYLVGTEPA++G+M+KY +P

Sbjct: 343 PIWPLIALSNIAQASAVVGIIIIISK-KQGERDISVPAAISAYLVGTEPMYGINLKYKFP 401

Query: 425 FVAGMIGSSVAGLLATTENNQANSIGVGGLPGFTLINVKYMGFFIOANVAIVVNFPLTY 484

++ MIGS +A + + V AN IGVGLPG ++I ++ + + M +AI+VP IT

Sbjct: 402 MLSAMIGSALAAAVCSAGVMANGIGVGGLPGILSIQPFWSIYLVNMLIAILVPAALTY 461

Query: 485 FFKK 488

K
Sbjct: 462 LMYK 465

An alignment of the GAS and GBS proteins is shown below:

Identities = 501/675 (74%), Positives = 573/675 (84%), Gaps = 2/675 (0%)

Query: 1 NMQFQGHAKALLEAIGGKENSIVTHCATMRMFVINDSSKAKVKVIEKLPSVKQTFPNAG 60
M ++ DAK+LL AIGGKENI VTHCATMRMFVIND++KA VK IB++ VKQTFPNAG
Sbjct: 1 NGKFEQDAKSLLT AIGGKENIKVTHCATMRMFVINDNNKANKKEIKSVVKQTFPNAG 60

Query: 61 QFQVIIGNDVPIFYNAFVAVSGIVGSKRAAKSAAQKNQFLQRVITMLAEIPTPIIPAI 120
QFQVIIGNDVP+FYN F AVS IRGVSKRAAKSAA+ NQN LQRV+TMLAEIPTPIIPAI
Sbjct: 61 QFQVIIGNDVPVFNDFPTAVSIEGVSKEAKSAAKSNQALQRVITMLAEIPTPIIPAI 120

Query: 121 IVGGLILGFRNILDVPPFELGQKVVGDVQVDSGGHPIWNTLVDVSTFMSGVDSFLALP 180
IVGGLILGFRNILL++VPFELGQ+V G D+G P+WNT+V VS FWSGV+ FLALP
Sbjct: 121 IVGGLILGFRNILESVPFELGQQVEKGLVFDAAQDPVWNTLVRVSPFWSGVNHLALP 180

Query: 181 GRAIFHFLPVGIVNSVTRKMGTTQILGIVLGICLVSPQLLNAYSVASTSAADIAKQWVN 240
GRAIFHFLPVGI NSVTRMGTTQILGIVLGICLVSPQLLNAY+VA T AA+IAQW N+
Sbjct: 181 GRAIFHFLPVGITSVTRKMGTTQILGIVLGICLVSPQLLNAYAVAGTPAAEIAKQWVMD 240

Query: 241 PGYFTVQKIGYQAQVIPALLAGLSLYLKIFWRKHIEPVVSMIFVFFLSLVPAILAHIV 300
PG+FT+ +IGYQAQVIPALLAGLSL+YLEIFWRK IPEVVSIMIFVFFLSL+PA+ILAHIV
Sbjct: 241 PGFPTINRIGYQAQVIPALLAGLSLAYLEIFWRGRIPEVVSIMIFVFFLSLIPAILAHIV 300

Query: 301 LGGPIGWTLGKWISAIVLIGLTGFVKMLFGAIFGALYAPFVITGLHMTNAIDTQLADTK 360
LGGPIGW+GK IS +VL GLTGFVKMLFGAIFGALYAP VITGLHMTNAIDTQLADT
Sbjct: 301 LGGPIGWTLGKGISFVVLGAGTGFVKMLFGAIFGALYAPLVTGLHMTNAIDTQLADTA 360

Query: 361 THTTGLNFMIALSNIAQSSAVLYYPMHRHDEKAQISLPAISAYLVTEPALFQVNVK 420
T TGLNFMIALSNIAQSSAV AYY M+RH+E+EA+ISLPAISAYLVTEPALFQVNVK
Sbjct: 361 TRITGLNFMIALSNIAQSSAVYALMNRHEEREAEISLPAISAYLVTEPALFQVNVK 420

Query: 421 YIYFPVAGMIGSGIAGLLATTFNVQANSIGVGLPGFLINVKMGYFFICMAVAIPIPL 480
Y+YFPVAGMIGS +AGLL+TFNVQANSIGVGLPGF++INVKIM FFICMAVAI +P+
Sbjct: 421 YVYFPVAGMIGSGIAGLLSTTFNVQANSIGVGLPGFMAINVKYIMFFPICMAVAIVVK 480

Query: 481 FLTLFFKSGILTKTEBKINVDIASTTETSKAREKAVVSGTKLSVSPLOGLAKPLD 540
FLT FF+KS I+TKTE+E +P+ +S +A K +GT +++ SPL+G K L
Sbjct: 481 FLTFVFRKSHIMTKTEBAKLFETPV-SDAVPATAPEK-TMQGTVITLTSPLTGEVKAIS 538

Query: 541 QASDPVFSQCLMGKGVVIDPSDGLVSPVDATVSVLPFTKHAIGLLTSBGVEVFLIIGND 600
+A DVPV+QG+MG+G ++ P+G LV+P DA VSVLPFTKHA I L+T+EG+E L+HIGND
Sbjct: 539 EAVDPVFAQGVNGQALLQPTBGVLVAPCAEVSGLVFFTKHAICLVTEBSLEMLIIGND 598

Query: 601 TVNMLCKGPTSHVAGQDTVKVQDKLITFDIEMIKRGGYIVSTPILITNQGEFRPEELIDL 660
TVNML+G+GP +V QGD VK G LI FDI I E GY ETP++TMQ F L
Sbjct: 599 TVNMLDGGQFEALVQDQGVKQQTILQFDIAISBAGYATSTETPLVNTNQDVPVTVESGL 658

Query: 661 PKQIGKQALVAVK 675
P+QIK L V A K
Sbjct: 659 PROIKVNDKLA VAVK 673

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 146

A DNA sequence (GBSx0152) was identified in *S. galactiae* <SEQ ID 487> which encodes the amino acid sequence <SEQ ID 488>. This protein is predicted to be dextran glucosidase DexS (treC). Analysis of this protein sequence reveals the following:

Possible site: 48

-222-

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

5 bacterial cytoplasm --- Certainty=0.3493 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

10 >GP:AAB65079 GB:U35633 dextran glucosidase DexS [Streptococcus suis]
 Identities = 383/547 (70%), Positives = 439/547 (80%), Gaps = 13/547 (2%)

Query: 1 MTIDKRKVVYQIYPKSYKDTTCNGVQDLRGIIRKLPYLAEGLIDMWLNPFYPSPORUNG 60
 Sbjct: 1 MTIDKRKVVYQIYPKSYKDTTCNGVQDLRGIIRKLPYLAEGLIDMWLNPFYPSPORUNG 60

15 Query: 61 YDISDYTAINPDFGTMDDFEEMIEVGROYRIDFMLDMVLNHCISIEHWFKALAGDRYYQ 120
 YDISDYTA+NPDPGTMDDFEEM+ VG++ I+PMLDMVLNHCIS+HSEW+KAL+GD+YYQ
 Sbjct: 61 YDISDYTAINPDFGTMDDFEEMVTVGKELGIEPMLDMVLNHCSTDEHWFQKALSGDQYYQ 120

20 Query: 121 DFFILRDNEPTDWSKPGGNAPFGDTGKYHLELFDITQADLNWRNADVRKELFKVNVFW 180
 DFFILRD PTDWSKPGGNAPFGDTGKYHLELFD+TQADLNWRN +R+ELFKVNVFW
 Sbjct: 121 DFFILRDQPTDWSKPGGNAPFGDTGKYHLELFDITQADLNWRNPHIREELFKVNVFW 180

25 Query: 181 KDKGVKGRFDVINLIGFDEILENCPIINDGKPAYTDRPITHDYLMNNAFSGQDSDFT 240
 +DKGVKGRFDVINLIGFDE E+CFINDGKPAYTDRPITHDYLMN+NNR+FG + FMT
 Sbjct: 181 KDKGVKGRFDVINLIGKDEAREDCPIINDGKPAYTDRPITHDYLMNNAFSGQDSDFT 240

Query: 241 VGEMSSATTIANCILYTAPEEREELGMAFNPHHLKVDYKDGQKWITINAFDPALRDLFHSWG 300
 VGEMSS+TTI NCILYTAPEER+ELGMAFNPHHLKVDYKDGQKWITIN FDF L+ LPH+WG
 30 Sbjct: 241 VGEMSSATTIANCILYTAPEEREELGMAFNPHHLKVDYKDGQKWITINAFDPALRDLFHSWG 300

Query: 301 EGMSEBGNWALFYNNHDOFALNRFDVVKFRNEGATMLAASIHLSRGTFYIYMGEEIG 360
 E M GNWGNWALFYNNHDOFALNRFD+V+ FR EGATMLAASIHLSRG
 35 Sbjct: 301 ERMSEBGNWALFYNNHDOFALNRFDVVENFRNEGATMLAASIHLSRGNNLTST----- 365

Query: 361 MLDPDYSSMDYVDIESLNAYQIMLDGKSGQREAFSIIIRAKSRDNRSPVMQWDDS----- 415
 + SS + + + + + S + R SR + P+
 40 Sbjct: 356 WVRSEYSSLTLTITTIATTTTWTWSLGMPTRCNWKVTRLR-PSKLSRSPVTIIPAPRCNGT 414

Query: 416 --TNAGFSBGAFLWKVGSYKEINVAKEKTLIFTFYQELIRLRQQLPIADGNYKAAFK 473
 T + PWLK GKSY+ INV +EKTG IFTFY+ LRK+LP+I++G+YKAA+K
 45 Sbjct: 415 LMTQASQQAFLWKAGKSYQITINVEQKTIPTFTFKGTHPLRKELPLISEGDIYKAAK 474

Query: 474 DNEKVYAFERHLDKEKLLVLNNFPFAEKVKIKLPENYLQQCVLLSNYKDTLTDEVTTLQPY 533
 D++KVYAFER L+ EKLLVLNNFPFAE++V++ L ++Y GQVL+SNY D L + + L+FY
 50 Sbjct: 475 DSQKVYAFERLNDKEKLLVLNNFPFAEVELDLADYAHGQVLISNYPDNKLGKIKLPY 534

Query: 534 QTLAILV 540
 Q LAI V
 50 Sbjct: 535 QALAIQV 541

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 489> which encodes the amino acid sequence <SEQ ID 490>. Analysis of this protein sequence reveals the following:

Possible site: 56

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

55 bacterial cytoplasm --- Certainty=0.3631 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 60 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 431/539 (79%), Positives = 486/539 (89%)

-223-

Query: 1 MTIDKKVVYQIYPKSYKDTGNGVGLGILGIEKPLAELGLDMVNLNPPYSPQRENG 60
 MTIDK+KVVYQIYPKSYKDTGNGVGL GII+KLPL ELGLDM+NLNPPYSPQRENG
 Sbjct: 1 MTIDKKVVYQIYPKSYKDTGNGVGLGILGIDKPLQLGLDMVNLNPPYSPQRENG 60

5 Query: 61 YDISDYTAIHPDGFIMDDFEREILVGRQYRIDFMLDMVLNHCSTIEHMFKKALAGDQRYQ 120
 YD+SDYTA+HPDGFIM DPE +++ ++L+ MLDMLNHCST+HEMF+KALAGD+YQ
 Sbjct: 61 YDVS DYTAIHPDGFIMDFENLVKAAKEHQIEMLDMVLNHCSTDEHMFQKALAGDPYQ 120

10 Query: 121 DFFILRDNPITDMSKPGGNWAPPGDGTGKYIHLFDITQADLANWKNADVRKELPKVNW 180
 DFFILRD PTDMSKPGGNWAPPGDGTGKYIHLFD+TQADLANWKN VR+EL KVNWF
 Sbjct: 121 DFFILRDQPTDMSKPGGNWAPPGDGTGKYIHLFDVYQADLANWKNFHVREELAKVNW 180

15 Query: 181 RDKGVKGRFDVIMILIGKDEILNCPINDGKPAYTDRPITHEDVLKMNASFGQDSDPMT 240
 RDKGVKGRFDVIMILIGKDE L +CP+NDGKPAYTDRPITH YL IN ASFGQDSDPMT
 Sbjct: 181 RDKGVKGRFDVIMILIGKDEELNDCPVNDGKPAYTDRPITHYIHLNQSFGQDSDPMT 240

20 Query: 241 VGEHSSTTIANCLLYTAPEREELSMAPNPHHLKVDYDKQKWTIMAFDPPALRDLFHWG 300
 VGEHS+TTI NC+LYTAPEREELSMAPNPHHLKVDY++GQRTIMAFD ALRDLFHWG
 Sbjct: 241 VGEHSATTIINCLLYTAPEREELSMAPNPHHLKVDYEWQKWTIMAFDFAALRDLFHWG 300

30 Query: 301 EGMSQNGWALFYTHBDQPALNRFDVVKRFNKGATMLAASIHLSRGTPYTYMGEEIG 360
 EGMS+NGWALFYTHBDQPALNRFDV FRNKGATMLAASIHLSRGTPYTYMGEEIG
 Sbjct: 301 EGMSQNGWALFYTHBDQPALNRFDVTFNFRNKGATMLAASIHLSRGTPYTYMGEEIG 360

35 Query: 361 MLDPOFYSMDYVDIESINAYQIMLDGKSGQEEAFSIRAKSRDNRSPVQNDSDTNAFG 420
 MLDPO+ SMDYVD+ESINAY +L GKS EEAF+II+AKSRDNR PVQND S +AGF
 Sbjct: 361 MLDPOFDSMDYVDIESINAYSSILVSGKSAEEAFIIXAKSRDNRKPVQNDSDSHAGF 420

40 Query: 421 SEGAPWLKVGKSYKEINVAKEKTLIPTFYQELIRLKLPLIADGNVYKAFKDKNKVYA 480
 + G PWL+VGKSY++INV EK G IF FYQ LI LRK+LEPIA+G+Y+AAFD++ VYA
 Sbjct: 421 TTGKFWLEVGSYRDNVETEKGRIPFPFYQRLIALRKELPIIAGDYRAAFKDSQAVYA 480

Query: 481 FERHLKKEKLLVNLNFFAEKVKIKLPENVLQGVLLSNVYKQVTLDETVTLPYQITLAIL 539
 FERHL +LVNLN+F+A++V++L P Y GQVL+SNY+ V++ E V L+PYQITLAIL
 Sbjct: 481 FERHLGDQCLLVNLNHFYADEVELELPPRYQHGQVLISNVEKVSICEKVLKPYQITLAIL 539

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 147

40 A DNA sequence (GBSx0153) was identified in *S.galactiae* <SEQ ID 491> which encodes the amino acid sequence <SEQ ID 492>. Analysis of this protein sequence reveals the following:

Possible site: 29
 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -3.03 Transmembrane 8 - 24 (8 - 25)

45 ----- Final Results -----
 bacterial membrane --- Certainty=0.2211 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

50

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 148

A DNA sequence (GBSx0154) was identified in *S. agalactiae* <SEQ ID 493> which encodes the amino acid sequence <SEQ ID 494>. Analysis of this protein sequence reveals the following:

Possible site: 57
 >>> Seems to have a cleavable N-term signal seq.
 ----- Final Results -----
 bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:BA03939 GB:AP001507 unknown conserved protein [Bacillus halodurans]
 Identities = 190/639 (29%), Positives = 331/639 (51%), Gaps = 34/639 (5%)
 Query: 6 TVVIMLVFLARKNLSLYELTVQTKFSIKVIEQINYINSFLAKNHLPAIAHSAGRYQLLG 65
 T ++ + AR L + ELT + S + + + NS+L + L A + + L +
 Sbjct: 8 TFLITQLLHARSYLPQELTQKLVSRPTVYNDLEKINSWLEEQGLKAV-KVRSQQLL 66
 Query: 66 DEKEHDKI---VSLLEAQFYLTQERVCVLIYLYSPCRREFVSNVHYQDFLKVSNVTLS 122
 DE+ ++I + L++ + + +ER + +Y RE + H D VS+NTT+
 Sbjct: 67 DERAKKEIPTKRLSLKSMHYESAQERKAWVYVYLTLRLEPLFLEHMDRTGVSRNTTID 126
 Query: 123 DIKMLRSKLAKRGISLTYTRAKGYSIAGDEMDKHQVAFQMITQLLE-----SPIGFW 174
 DIK L+ +L ++L + R GY++ GDE DK + ++Q L SPI +
 Sbjct: 127 DIKCLDELNPNHFALEFERKDGTYTISGDETDKRALVYVLSQALPQQNWETELSPRIF 186
 Query: 175 SINVILSSWKFALSYEKLEKTVFYFESFOLSPIQ---DRLEKSLYFIILLICRYQSVSD 231
 + F + E+L+K + ES ++ IQ D L +L + R +
 Sbjct: 187 LRTKRDNGRIFTI--BELQKVYDVISESEKVLKICYTDDVLHLSLFLLEPMKRVAKG-- 242
 Query: 232 RVLQGSPIVSEQLK----ELATTIVTNLSQDISLSPKLDQKEKDYITLLISGCF---- 281
 ++ P+ + LK E ++ L Q + P D++ T ILS
 Sbjct: 243 KFIKVPFLKQVLKGTKEYEAAKVMSPKLEQAQGVHY-PDEEVLFLTHILLSSKINYANG 301
 Query: 282 EGEQTKDDDFEALAKAIVDEMETSVLNFSNKEELLQGLRHIIIPAYFLKYGITLSDG 341
 E E K+ + ++V++ + + F KE L + L HI PA++R+KYGL ++
 Sbjct: 302 EIPSRKESQLTHIVTSMVNDPQKYACVFFEREKELLEKNLFFHIKPAFYRIKYGLEVENN 361
 Query: 342 YTONIKEHYSDLFLLVKKALRPLEEQVGL-IPDSEISYFVIHFQGYLRQSGGTQMSYKA 400
 ++IK Y +LFL +K + LE VG + D+E+++ +HF G++R+ G + KA
 Sbjct: 362 IAESIKTYPPELFLTRKVVHYLRYVGSVDNDEVAFTIMHFVGWRRREGTIPTKRRKA 421
 Query: 401 LILCPNGVSSSLVIKEKRLGLFPQIHFRVSKIEBQLKLDNQTVMVSTIFVETKKPNY 460
 LI+C NGV +S +K +L GLFP + I + + + ++T E P +
 Sbjct: 422 LIVCANOVGTSQFLNQLLEGLFPAVDIITKTSIREYEKTFVPEVDITSTTSIFENGVPIF 481
 Query: 461 LVSIAMT-ABQVQQLKELVISDFPKACLDPPOLDQLIATIKKYAHVHCERELKIALRTMV 519
 +V+ ++T E+ + LK ++ + + ++ L IK++ +V E+ L LR
 Sbjct: 482 IVPNLTETRKERLLKSVHVALDELGAQKYSIEGLMDVIRKGNVDDEKALYQDLRRFF 541
 Query: 520 KQD--ILRKDVRLIAQLITESTYQTSBQNMWKEAIRLAAPLLASGCITESYPEANIE 577
 Q I K +P L+QL+T+ Q + +W+RAI+LAAPLL G +TESY + NI+
 Sbjct: 542 TQPTFTPGKQKPDNLQLLTEDIQRLQRBQVTHQBAIQAAKPLLLKGNVSTSVYKKMIK 601
 Query: 578 KVEEFGPFINLKGIAIPHARPEQGVNSVGMMLVLEQ 616
 +E+FGP++ + ALPHA+PEQGV +GMS+L L++P
 Sbjct: 602 NIKRQFYMIILAPHPAIPHAKPEDGVQQLGMSIIMLKKP 640

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 495> which encodes the amino acid sequence <SEQ ID 496>. Analysis of this protein sequence reveals the following:

Possible site: 57 or 61
 >>> Seems to have no N-terminal signal sequence

-225-

INTEGRAL Likelihood = -0.64 Transmembrane 123 - 139 (123 - 139)

----- Final Results -----

- 5 bacterial membrane --- Certainty=0.1256(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

- Identities = 187/624 (29%), Positives = 327/624 (51%), Gaps = 20/624 (3%)
- 10 Query: 1 MVDNKTIVIMLFLARKNLSLYELTVQTKPSIKVIIIBQINYLNSFLAKNHLPAIAHSAGR 60
 M+ ++ + + +F K SL K S + I+ I +N L+ LF IA
 Sbjct: 35 MLSHELIRNYQLFSKYKGHSILKAPESILKASKRHIIADIAKINDTISLYQLFLIALDR-- 92
- 15 Query: 61 YQLL--GDEKEHDKIVSLLEAEQFYLTQEERVCILYISFCRRFVSNNVHYQDFLKVSKN 118
 QL+ D E D + +L YL Q+ER+ +I +Y +EF+S H + L++S+N
 Sbjct: 93 -QLVYPPDLTEKDLLRMFLPTLDLQFQDERLMDIIITYIMAKFISINHLESLLRLSRN 151
- 20 Query: 119 TTLSIDIKMRSLKLRGISLITYTRAKQYSLVGDEMDEHQVAFQMITOLLESPIGFHSIN 178
 + ++D+ ++R ++ ++D Y R GY G+ + + + LL+ G W +Y
 Sbjct: 152 SVIADINLVRDRVQAFQVTLAYNRQDGYFFBGEFLARLLESASVSSLLQVTSQGMVFSY 211
- 25 Query: 179 ILSNKKFALSIEKLEKTVFYFSPQLSPIDRLEKSNIFYLITLCR-YQRSDV-RVLQ 236
 +L + + T+E L+ I ++L +YF L+ R +R+V
 Sbjct: 212 LLEHGLFDQKQKVMATLELSRENHLTFISEKRLDIYFFCLAHRRPFRHVRASAVDT 271
- 30 Query: 237 SPIVSEQLKELTTIIVTNLSQDISKLPLDQEKDYITILSGCFEG--BGTKDDDFEA 294
 P+ S ++ + ++ N P +EK + L GC +G E ++
 Sbjct: 272 FLASPAVETMVDQLLVNF-----PSLTBEKYLVSRLGCTIGDLELVFQQPYDI 323
- 35 Query: 295 LKAIVDEMETSLLMFNSKEELLQGLRHIIIPAYFLAYGLTGDGGYTNKIKHYSDLF 354
 + + I++ + + L+ ++ EL Q L R++PAY+L Y + + + IK+Y LF
 Sbjct: 324 NEE-IINSVAVNTGLSITDITPELRQNLVSHLLPAYTFLYDINLTNPLKQIKQGVESLF 382
- 40 Query: 355 LLWKKALRPLEQVGL-IPDSEISYFVIHPGGLRQSGGTQSMYSKALICPNVSSSLV 413
 LNK++L ELE+Q+G + + E++YF IHFG +L+ S AL +CPNG+SSSL+
 Sbjct: 383 YLVKRSLSPLKQLKQSVNEDRVAYFTIHFGNQLQPKKPSNQLVALSVCPNIGSSSLN 442
- 45 Query: 414 IKEKLGFLFPQIHFRVSKIEQLKIDNQTDMVESTIFVETKKNYLVSLMTAQVQQ 473
 ++ L+ LFFQ+ F R+ ++++KL+D ++D++EST+ + KP Y+ +M +
 Sbjct: 443 LEATLKELFQLQFIRIHQLDKIKLDPASFLIFSTVAFDCAKPVTVTQALMGFVEH 502
- 50 Query: 474 LKELVSDFFKACLDLPQLDQIATIKKYAHVHCBEELKAL-RIMVQDPIRKQVRELL 532
 LK++V DF + F LD L++ I K+ + +E L L R ++ + L
 Sbjct: 503 LKRWCDFFHLFLSEQFALDGLSLTIHKHTTTNKGKLVSDLSRYLIGNHLTEKGGGL 562
- Query: 533 HQLITERTYQTSSEQMMKCAIRIAKPLASKITESPEAMIEKVERGGPFINLKGII 592
 L+T + + + +N+EAIRLAA+FL I SY + MI+ V E G + I L +
 Sbjct: 563 LDLTLADFIQADAVSDWQEIRLAAQFLLEKQMIETSYIDGMIDSVNELGAYIVLAFKV 622
- Query: 593 AIHPARPEDGUNSVMGMSLVLEQP 616
 A+PHA PE G +GMS+L L++E
 Sbjct: 623 AVHAAPEKGTQLGMSLLCLKPE 646

- 55 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 149

A DNA sequence (GBSx0155) was identified in *Sagalactiae* <SEQ ID 497> which encodes the amino acid sequence <SEQ ID 498>. Analysis of this protein sequence reveals the following:

- 60 Possible site: 22
 >>> Seems to have no N-terminal signal sequence

-226-

```

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3665(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 499> which encodes the amino acid sequence <SEQ ID 500>. Analysis of this protein sequence reveals the following:

```

Possible site: 22
>>> Seems to have no N-terminal signal sequence

```

```

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3665(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

An alignment of the GAS and GBS proteins is shown below:

```

Identities = 33/35 (94%), Positives = 35/35 (99%)

```

```

Query: 1 MEKEAKQIIDLKRNLFKIDVRAQKDEEKVFMRTAW 35
+EKEAKQ+IDLKRNLFKIDVRAQKDEEKVFMRTAW
Sbjct: 1 LEKEAKQMIDLKRNLFKIDVRAQKDEEKVFMRTAW 35

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 150

A repeated DNA sequence (GBSx0156) was identified in *S.agalactiae* <SEQ ID 501> which encodes the amino acid sequence <SEQ ID 502>. This protein is predicted to be a repeat-associated protein in rhsc-phrb intergenic region. Analysis of this protein sequence reveals the following:

```

Possible site: 44
>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood = -4.57 Transmembrane 29 - 45 ( 28 - 43)

```

```

----- Final Results -----
bacterial membrane --- Certainty=0.2826(Affirmative) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

A closely-related DNA sequence was identified in *S.agalactiae* <SEQ ID 1035> which encodes the amino acid sequence <SEQ ID 1036>. Further related GBS sequences are: <SEQ ID 9067>, <SEQ ID 9068>, <SEQ ID 9497>, <SEQ ID 9498>, <SEQ ID 9733>, <SEQ ID 9734>

A related repeated DNA sequence was identified in *S.pyogenes* <SEQ ID 503> which encodes the amino acid sequence <SEQ ID 504>. Analysis of this protein sequence reveals the following:

```

Possible site: 44
>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood = -4.57 Transmembrane 29 - 45 ( 28 - 48)

```

```

----- Final Results -----
bacterial membrane --- Certainty=0.2826(Affirmative) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

A related GBS gene <SEQ ID 8547> and protein <SEQ ID 8548> were also identified. Analysis of this protein sequence reveals the following:

```

5      Lipop Possible site: -1      Crend: 5
      Mcg: Discrim Score:      -7.73
      GvK: Signal Score (-7.5): -3.88
      Possible site: 44
  >>> Seems to have no N-terminal signal sequence
      ALOM program count: 1 value: -4.57 threshold: 0.0
10     INTEGRAL Likelihood = -4.57 Transmembrane 26 - 42 ( 25 - 45)
      PERIPHERAL Likelihood = 2.12 334
      modified ALOM score: 1.41

      *** Reasoning Step: 3

15     ----- Final Results -----
           bacterial membrane --- Certainty=0.2826 (Affirmative) < succ>
           bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
           bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

20 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 7071> which encodes the amino acid sequence <SEQ ID 7072>. An alignment of the GAS and GBS sequences follows:

```

      Score = 767 bits (1960), Expect = 0.0
      Identities = 375/377 (99%), Positives = 375/377 (99%)

25     Query: 4 MIDFIISIDDCAVELDSRQGWKIRSPILSTLLFLVFCVQLAGIETWKEMDFIENNEPLFA 63
      MIDFIISIDDCAVELDSRQGWKIR PLSITLLFLVFCVQLAGIETWKEMDFIENNEPLFA
      Sbjct: 1 MIDFIISIDDCAVELDSRQGWKIRYPLSTLLFLVFCVQLAGIETWKEMDFIENNEPLFA 60

30     Query: 64 TYVDLSSEGCSSHDTLERSVLVNSDRLEKELKVQFBQSLTSLDAVHQLISVDGKTIIRNGR 123
      TYVDLSSEGC SHDTLERSVLVNSDRLEKELKVQFBQSLTSLDAVHQLISVDGKTIIRNGR
      Sbjct: 61 TYVDLSSEGCSSHDTLERSVLVNSDRLEKELKVQFBQSLTSLDAVHQLISVDGKTIIRNGR 120

35     Query: 124 KNQKPVHIVTAYDGGHLSLQGVAVEEKSNEIVAIPQLLRITIDIRKSIIVTIDAMGTQTAI 183
      KNQKPVHIVTAYDGGHLSLQGVAVEEKSNEIVAIPQLLRITIDIRKSIIVTIDAMGTQTAI
      Sbjct: 121 KNQKPVHIVTAYDGGHLSLQGVAVEEKSNEIVAIPQLLRITIDIRKSIIVTIDAMGTQTAI 180

      Query: 184 VDTIIGKADYCLAVRGNEQETLYDDIALYPSDVNLLBELQENAOYYQTVESRSGQIEVRE 243
      VDTIIGKADYCLAVRGNEQETLYDDIALYPSDVNLLBELQENAOYYQTVESRSGQIEVRE
      Sbjct: 181 VDTIIGKADYCLAVRGNEQETLYDDIALYPSDVNLLBELQENAOYYQTVESRSGQIEVRE 240

40     Query: 244 YWVSDIKWLQCNHPKWKHLRGIGMTRNTIDKDGQLSQENRYFISFKPDVLTFCANVRG 303
      YWVSDIKWLQCNHPKWKHLRGIGMTRNTIDKDGQLSQENRYFISFKPDVLTFCANVRG
      Sbjct: 241 YWVSDIKWLQCNHPKWKHLRGIGMTRNTIDKDGQLSQENRYFISFKPDVLTFCANVRG 300

45     Query: 304 HWQIBSMHWLLDVVYHEDHHQTLDKRAAFNLNLRIMCLYFLKVMVFPKDLISYRKQRY 363
      HWQIBSMHWLLDVVYHEDHHQTLDKRAAFNLNLRIMCLYFLKVMVFPKDLISYRKQRY
      Sbjct: 301 HWQIBSMHWLLDVVYHEDHHQTLDKRAAFNLNLRIMCLYFLKVMVFPKDLISYRKQRY 360

50     Query: 364 ISVHLEDYLVLQFGERG 380
      ISVHLEDYLVLQFGERG
      Sbjct: 361 ISVHLEDYLVLQFGERG 377

```

A further related DNA sequence was identified in *S.pyogenes* <SEQ ID 9087> which encodes the amino acid sequence <SEQ ID 9088>. A further related DNA sequence was identified in *S.pyogenes* <SEQ ID 9089> which encodes the amino acid sequence <SEQ ID 9090>. The GAS and GBS proteins are 100% identical.

There is also homology to SEQ IDs 7018 and 8548.

SEQ ID 8548 (GBS318) was expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 46 (lane 5; MW 70kDa).

GBS318-GST was purified as shown in Figure 203, lane 3.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 151

- 5 A DNA sequence (GBSx0157) was identified in *S.agalactiae* <SEQ ID 505> which encodes the amino acid sequence <SEQ ID 506>. Analysis of this protein sequence reveals the following:

Possible site: 34
>>> Seems to have an uncleavable N-term signal seq

- 10 ----- Final Results -----
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

- 15 The protein has no significant homology with any sequences in the GENPEPT database, but there is homology to SEQ ID 496.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 152

- 20 A repeated DNA sequence (GBSx0158) was identified in *S.agalactiae* <SEQ ID 507> which encodes the amino acid sequence <SEQ ID 508>. Analysis of this protein sequence reveals the following:

Possible site: 48
>>> Seems to have no N-terminal signal sequence

- 25 ----- Final Results -----
bacterial cytoplasm --- Certainty=0.1054(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>

- 30 The protein has homology with the following sequences in the GENPEPT database:

>GP:BAB03941 GB:AP001507 unknown conserved protein [Bacillus halodurans]
Identities = 26/82 (31%), Positives = 52/82 (62%), Gaps = 2/82 (2%)

- Query: 2 LRIGTACSGGLGSSPMVQWNIESTLKDLGVSDVEVHYDLGGADPSAADVWIVGRDLES 61
+I C3 G G+S ++W+E+L LG++ +V+ D+ A +D I +L +S
35 Sbjct: 1 MKILVCGLCQCTSLIKQVETVLSQLGIA-ADVENTDVSSASSEQSDFIITSKELAS 59

Query: 62 -AGHLGDVRTILNIIIDMELEK 82
A H + I+N+ DM+E++

- 40 Sbjct: 60 LASHPSKIVIVNNYFDMSEIKQ 81

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 509> which encodes the amino acid sequence <SEQ ID 510>. Analysis of this protein sequence reveals the following:

Possible site: 49
>>> Seems to have an uncleavable N-term signal seq

- Final Results -----
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
50 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

-229-

Identities = 27/90 (30%), Positives = 51/90 (56%), Gaps = 1/90 (1%)

Query: 1 MLPITGACSGGLGSSFMVQMNITSLTKDLGVSDVVEHYDLGADPSAADVWLVGRDGLD 60
 N+I T CG+G+GSS +++M +E+I LG+ DV+ E D A AD+++ ++ +D
 5 Sbjct: 8 MIKIVTQENIGSIGSLLLRMKVERIASSLGI-DVEABS CDSNAVVGKADLPFTVKEFKD 66

Query: 61 SAGHLGDVRIINSLIIMDELRLRLVTCIQE 90
 V I+ S + ++ E + + +E
 10 Sbjct: 67 IPPEDAKVCIVKSYTNKKKIERDLVPVKK 96

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 153

A DNA sequence (GBSx0159) was identified in *S. agalactiae* <SEQ ID 511> which encodes the amino acid sequence <SEQ ID 512>. Analysis of this protein sequence reveals the following:

Possible site: 20
 >>> Seems to have an uncleavable N-term signal seq

----- Final Results -----

bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S. pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 154

A DNA sequence (GBSx0160) was identified in *S. agalactiae* <SEQ ID 513> which encodes the amino acid sequence <SEQ ID 514>. This protein is predicted to be *sgaT*. Analysis of this protein sequence reveals the following:

Possible site: 16

>>> Seems to have a cleavable N-term signal seq.

INTEGRAL Likelihood = -14.97 Transmembrane 424 - 440 (411 - 447)
 INTEGRAL Likelihood = -8.86 Transmembrane 224 - 240 (221 - 248)
 35 INTEGRAL Likelihood = -7.27 Transmembrane 134 - 150 (124 - 167)
 INTEGRAL Likelihood = -7.11 Transmembrane 321 - 337 (314 - 349)
 INTEGRAL Likelihood = -6.64 Transmembrane 379 - 395 (370 - 397)
 INTEGRAL Likelihood = -6.21 Transmembrane 96 - 112 (94 - 115)
 40 INTEGRAL Likelihood = -6.05 Transmembrane 267 - 283 (257 - 289)
 INTEGRAL Likelihood = -3.13 Transmembrane 18 - 34 (17 - 35)
 INTEGRAL Likelihood = -2.55 Transmembrane 151 - 167 (151 - 167)
 INTEGRAL Likelihood = -0.32 Transmembrane 42 - 58 (42 - 58)

----- Final Results -----

bacterial membrane --- Certainty=0.6986(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAB52363 GB:AL109747 putative integral membrane protein
 [Streptomyces coelicolor A3(2)]
 Identities = 202/453 (44%), Positives = 292/453 (63%), Gaps = 22/453 (4%)

-230-

Query: 7 FLVN-IASTPAILVALLAIGLVIRKRGVDPVIGKGIKTFVGVFLVSGGTGIVQNSINPF 65
 FLVN I S PA L+ +I +GI KK V V G I K +G L+V G G+V +SL+P
 Sbjct: 10 FLVNSILSQPAYLIGIITAVGLAALKKSVGQTVGGNATKATIGLLLVGGAGLVSSSLDPL 69

5 Query: 66 GKMFHFAHLVGVVNNNAIVAVALTQYGSNTALIMLAGHIFMLIARPTKFKYIFLTGH 125
 G+M + GV+P NEAIV +A +++G+ A +H+ G + ++ ARPT +Y+FLTGH
 Sbjct: 70 GRMIQGTGTGTHGVIPINBAIVGLAQSEFGARVAVMLILGFLVSLALARPTFLRYVFLTGH 129

10 Query: 126 HTLVMAACMIAVIFAVAGFTSFLLIFGGLALGIMSVSPAFVQKMIQLTQNDKVALPH 185
 H L+MA + ++ A AG S +++L GG+ +GI++ DAF + ++TQND +A+GHF
 Sbjct: 130 HMLFATMLITIVMATAGQSSVAVVLCQGVILGILLVALPAFAHFWPKYKTCNDTLAIGHF 189

15 Query: 186 GSLGTYLSSGFIGGIVGDKSKSTEDIKPKKSLSPLRDSTVITISIMAILIYLIVAV- 239
 G+ GY +SG G +VG S+STE++K P+ L PLRDS V+ +SM +IY+ +++
 Sbjct: 190 GTAGYIVSGATGQLVGGKSKSTEMKLPGLRPLRDSMVATAGSMVLIYVMSLLFLAKV 249

20 Query: 240 -----PAGEAYIAKEISNVAKGLVYALQAGQPAAGVFVILAGVRLILGEIVPAFG 291
 FAG ++ N L+ ++ QF GV VIL GVR ILGE+VPAFG+
 Sbjct: 250 GQDAAPKAPAGSG--GDPADVGNVIMQSMQGLQFGIGVAVILPGVRIITELGVPAFG 307

25 Query: 292 ISEKLVNSKPAKDCTPIVYPANAVLIGFISFVGGVLSVMIMI-----VTGTIVILPG 346
 I+ ++VD +KPAD DIV+FYA NAVLIGFI SF+GGL + +I G ++LFG
 Sbjct: 308 IAGRVVPKAPKALDAPIVFPYANAVLIGFIPFGLTGLAALIVPFPAGLALVILPG 367

30 Query: 347 VVPHFFQGTAGTVIGNAGSGVGRGATGAPVQGLISFLPLFPLMPVGLGFGKSTSPSAD 406
 +VPHFF G AGV GNA+GG RGA +G+P+ G+LI+PLP L+ LG G +TF DAD
 Sbjct: 368 LVPHFFTGAGVGYGNATGGRGGAAGVSGFLNGLITPLPAILLKLALGSGEANTTFGDAD 427

Query: 407 PGLTGIIIGALNIVGGAIAIVIGIVVILIGLPG 439
 PG G +LG++ +G ++ ++ L+ L G
 Sbjct: 428 PGWFGVLSIGKLDGTAGLIGMLIFGLLILAG 460

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 515> which encodes the amino acid sequence <SEQ ID 516>. Analysis of this protein sequence reveals the following:

35 Possible site: 34
 >>> Seems to have a cleavable N-term signal seq.

INTEGRAL	Likelihood = -0.33	Transmembrane	330 - 346 (315 - 353)
INTEGRAL	Likelihood = -0.17	Transmembrane	227 - 243 (221 - 246)
INTEGRAL	Likelihood = -4.62	Transmembrane	127 - 143 (126 - 145)
INTEGRAL	Likelihood = -4.25	Transmembrane	269 - 285 (266 - 291)
INTEGRAL	Likelihood = -3.77	Transmembrane	43 - 59 (41 - 62)
INTEGRAL	Likelihood = -3.66	Transmembrane	98 - 114 (91 - 116)
INTEGRAL	Likelihood = -2.76	Transmembrane	146 - 162 (145 - 163)
INTEGRAL	Likelihood = -1.59	Transmembrane	308 - 324 (308 - 324)

45 ----- Final Results -----
 bacterial membrane --- Certainty=0.4333 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP:CB52363 GB:AL109747 putative integral membrane protein
 [Streptomyces coelicolor A3(2)]
 Identities = 162/387 (41%), Positives = 245/387 (62%), Gaps = 17/387 (4%)

55 Query: 8 IRDILKEPAPLGLIAPAGLVALKTPAHKVLITGLPILGYIMIAVAGGVIVNLDPLAK 67
 +IL +PA+L+G+I GL ALK + + G + LG L++ AGAG++ ++LDEL +
 Sbjct: 12 VNEILSQPAYLIGIITAVGLAALKKSVGQTVGGATKATIGLLLVGGAGLVSSSLDPLGR 71

60 Query: 68 LIEHGSFISGVVNNNAIVTSVAQKILGVETMSILVGLLNLINAFARPTKFKYIFLTGHHS 127
 +I+ GV+P NEA+ +AQ G ++++G L+L+LA ARPT +Y+FLTGH
 Sbjct: 72 MIQGTGTGTHGVIPINBAIVGLAQSEFGARVAVMLILGFLVSLALARPTFLRYVFLTGHM 131

Query: 128 FFMACLLSAVLGAVGFKGSLIIL-DGFLLGANSALSPAIGQYTLKVTDGDIAMGHFG 186
 FMA LL+ V+ G +GS+ ++L G L+G PA +T KVT D +A+GHFG

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Sbjct: 132 LFMATLLTIVMATAG-QSSVAVVLGGVVLGILLVALEFAHPWTKVTGNDTLAIGHFG 190
 Query: 187 SLGYLSAWVGSKVGKDSKDTEDLQISEKWSFLRNTTISTGLIMVIFYLAT--VASVL 243
 + GY +S G V GK+S+ TE++ + B FLR++ ++T L MV+ YLV + +A V
 5 Sbjct: 191 TAGYIVSGATGQLVGNKRSSTREMKLPGLRFLRDSMVATALSMLLYLVMSLLFLAKVG 250
 Query: 244 RNASVAEELAAQGNP-----PIFAIKSGLTFAGVVAIVYAGVRMILADLIPAFQGIAN 296
 ++A+ +G +P + + GL F +GVA++ GVR IL +L+PAFQGI
 10 Sbjct: 251 QDAAPKAFAGSGGDPADAVGNVLAQSVNQLGRLGIGVAVILPGVKTILGELVFAFGGLAG 310
 Query: 297 KLIPNAIPAVDCAVFFPYAPTAVIIGFASSFVGELLMIL----GVAGGLVLIIGMVP 351
 ++P A PA+D + FPYA AV+IGF SF+GEL G L G L++PG+VP
 Sbjct: 311 RVVGAKPALDAPIVFPYTAQNAVLIQIFISFLGGLTGLAALIWVNFAPGLALVLPGLVP 370
 15 Query: 352 HFFCGATAEIFGNSTGGRGAMIGASL 378
 HFF G A ++GN+TGRRGA +G+ L
 Sbjct: 371 HFTTGAAGVYGNATGGRGAAVGSFL 397

An alignment of the GAS and GBS proteins is shown below:

20 Identities = 174/376 (46%), Positives = 258/376 (68%), Gaps = 2/376 (0%)
 Query: 1 MKGLLOFLVNIASTPAILVALIAIIGLVQKKGVPDIVKGIKTFVGLVVSQGTIVQN 60
 M+ LL P+ +I PA L+ LIA GLV K ++ G + +G+L++ G G+
 25 Sbjct: 1 MEALLSFIRDIIEKPAFLMGLIAFAGLVALKTPAHKVLVTGLTLPILGYLMVAGAGVTV 60
 Query: 61 SINPFGKINFEHAFHLGVVFNNEAIVAVALTICYGSATALIMLAGMIFNIIARETFPKFYI 120
 +L+P K + EH F + GVVFNNEA++VA G T I++ G++ N+ ARFT+FKYI
 Sbjct: 61 NLDPLAKLIEHGFSTITGVVFNNEAVTSVAQLGLVETMSILVGLLLNLAFARFTRFKYI 120
 30 Query: 121 FLTGHHTLYMACMIAVIFAVAGFTSPSLILPGGLALGIIMSVSPAFVQKYMILQTNQKV 180
 FLTQHH+ +MAC++ + GF LI+ G LG ++SPA Q+Y +++T D+
 Sbjct: 121 FLTGHHSFPMACLLSAVLGA VGFKGSLIILDGFLLGANSALSPAIGQQYTLKVTGDDEI 180
 35 Query: 181 ALGHFGSLGYLSQFIGGIVGDKSKSTEDIKFPKSLFLRDSVTSITISMAIYLI--VA 238
 A+GHFGSLGY+LS ++G VG SK TED+ + SFLR++T+S + M I YL+ VA
 Sbjct: 181 AMGHFGSLGYLSAWVGSKVGKDSKDTEDLQISEKWSFLRNTTISTGLIMVIFYLATVA 240
 Query: 239 VFAGEAYIAKEISNGVNLVYALQLAQFAAGVFVILAGVRLILGEIVFAFGI3EKLVP 298
 A +A+E++ G N ++A++ FA GV ++ AGVR+IL +++PAF+GI+ KI+P
 40 Sbjct: 241 SVLRNASVAEELAAQGNPPIFAIKSGLTFAGVVAIVYAGVRMILADLIPAFQGIANKLIP 300
 Query: 299 NSKPALDCPIVYPYAPNAVLIIGTISFVGGLVSKIMVIMVTGTTVLPGVVFHFFCGATAG 358
 N+ PA+DC + +PYAP AV+IGF SSFVGL+ M++ V G +I+PG+VFHFFCGATA
 Sbjct: 301 NAIPAVDCAVFFPYAPTAVIIGFASSFVGGLLMILGVAAGVLIIPGMVVFHFFCGATA 360
 45 Query: 359 VTGNASGGVVGATIGA 374
 + GN++GG RGA IGA
 Sbjct: 361 IFGNSTGGRGAMIGA 376

50 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 155

A DNA sequence (GBSx0161) was identified in *S.agalactiae* <SEQ ID 517> which encodes the amino acid sequence <SEQ ID 518>. This protein is predicted to be transketolase, N-terminal subunit (tkt). Analysis of
 55 this protein sequence reveals the following:

Possible site: 45
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

60 bacterial cytoplasm --- Certainty=0.3680 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

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bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

5 >GP:AA98676 GB:U67515 transketolase [Methanococcus jannaschii]
 Identities = 106/269 (39%), Positives = 158/269 (58%), Gaps = 4/269 (1%)

Query: 11 LRKFATETIRLNTLETIANHLGPHGYGSSLSIVSALAVLYGDMIDINPKPKESDRYVMS 70
 L + A + R N + + + GH GSSLS + + LZ +M+ +P+ + DRD VLS
 10 Sbjct: 10 LEKIAKKVRYNIVOMVLGAKSGHPGSSLSATDITVALYFKIMNYSPPNPFYKDRDRFVLS 69

Query: 71 KGHAGPALYSTLYLKGFFDKTFLHSLNNTNGTKLPSHPDRNLTGPDIVTGSLLQGISIAT 130
 KSHA PADY+ L G ++ L L KL HP + TPG+++ TGSLLQ3 S A
 Sbjct: 70 KGHAAAPLAVLSLSLGIIEEELWKLRLEGKLGQHPMD-TPGVICTGSLGQGSFAAV 128

15 Query: 131 GIATYAKTIENSSYYTIVTVDGELNBSQCWEAIQFAAHQLHLHIVFDVDMNKQDLGLTA 190
 G+A +++ + Y Y ++GDGE EG WEA AAH++L +LI F+D NK Q+DG T
 Sbjct: 129 GMAIGCRDLKLNNTVTVLLGDGECQBGIVWEAAMAAHYKLDNLAFIDRNKLIQDQCTE 188

20 Query: 191 DICNPGDFVAKFEAFGDFAVRVKGGDDIRADKAITFQDSNVRPKCIVLDSIKGQGVKE 250
 D+ + GD ARFEAFG+D + G + E I ++ + + +FK I+ ++KG+G
 Sbjct: 189 DVMSLGDITAKFEAFGDFVFEIDGHNFEELNTEKAKSMKNKPKPMIAYTVKGGVSP 248

Query: 251 LEEELASNHSLRFDLQCKTMLERALISLE 279
 +E + H E+ +Q L++L L E
 25 Sbjct: 249 MENNVAFHGFAPNEEQ---LQALEELSE 274

A related DNA sequence was identified in *Spyogenes* <SEQ ID 519> which encodes the amino acid sequence <SEQ ID 520>. Analysis of this protein sequence reveals the following:

Possible site: 26
 30 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -0.75 Transmembrane 58 - 74 (57 - 74)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.1298(Affirmative) < succ>
 35 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

A related sequence was also identified in GAS <SEQ ID 9165> which encodes the amino acid sequence <SEQ ID 9166>. Analysis of this protein sequence reveals the following:

Possible site: 54
 40 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -0.75 Transmembrane 40 - 56 (39 - 56)
 ----- Final Results -----
 45 bacterial membrane --- Certainty=0.130(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

50 Identities = 82/246 (33%), Positives = 129/246 (52%), Gaps = 15/246 (6%)

Query: 18 IRNFTLETIANHLGPHGYGSSLSIVSALAVLYGDMIDINPEKFE-SDRDYVLSKGHAGP 76
 +R +++ + GH G + VLA+ M+INP+ + S+RD +LS GH
 55 Sbjct: 82 VRTLSMDAIQANGSHGFLPMGAAPMAYVLNHPFMNINPKTSRNSNRDRPILSAGHGA 141

Query: 77 ALYSTLYLKGFFDKTFLHSLNNTNGTKLPSHPDRNLTGPDIVTGSLLQGISIATGIAYA 135
 LYS L+L G+ L + G+K P HP+ N T G+ + TTG LQGGI+ A G+A A
 Sbjct: 142 MLYSLHLAGYDLSVEELKNFRQNGSKTPGHPVENHTGVEATTGFLQGISIANAVGMA 201

60 Query: 136 QK-----IENSSYYTIVTVDGELNBSQCWEAIQFAAHQLHLHIVFDVDMNKQDL 185
 + + +YT+ + GDG+L EG EA A R +L I++ D N L
 Sbjct: 202 EAHLAAPFKPGFDIVDHYTFALNGDGLMBSGVSQAASAGHLLKELKILVLYDSNDISL 261

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Query: 186 DGLTADICNPGDFVAKPEAPGFDAVRVK-GDIEAIDKAIKTFQDSNSVRPKCIVLDSIK 244
 DG T+ + D +FEA+G+ +VK G+D+E I AI+ + + +P I + +I
 Sbjct: 262 DGPTS-MAPTEDVKGRPEAYGQHILVKDGNLDEITAAIEAAK-AETEKPTIEIVKTI 319

Query: 245 GGGVKE 250
 G G ++
 Sbjct: 320 GFGAEK 325

- 10 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 156

A DNA sequence (GBSx0162) was identified in *S. galactiae* <SEQ ID 521> which encodes the amino acid sequence <SEQ ID 522>. Analysis of this protein sequence reveals the following:

- 15 Possible site: 43
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.27 Transmembrane 53 - 69 (53 - 69)
- 20 ----- Final Results -----
 bacterial membrane --- Certainty=0.1107 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

- 25 A related GBS nucleic acid sequence <SEQ ID 9499> which encodes amino acid sequence <SEQ ID 9500> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

- >GP:AB98674 GB:U57515 transketolase'' [Methanococcus jannaschii]
 Identities = 100/301 (33%), Positives = 171/301 (56%), Gaps = 7/301 (2%)
- 30 Query: 6 KEMRLVYRDFLLQANQENKQITVLEADLSSSMSTNALASEFGPKRYINGLIMEAEMVGLAA 65
 K MR Y + L++ ++ + + VL+ADLS S T A EF +R+ N G+ E M+GAA
 Sbjct: 9 KGMRRKYGETLIELGKKYENLVILDADLSGSTQTAMFAKEPPEPFNAGVABQNMIGMAA 68
- 35 Query: 66 GLAIKGYKPYLHTPGPFASRRVFDQVPLSLGYSCLSATIIGSDAGISABMNGGTHMPFE 125
 GLA G + +P FAS R ++ + + Y +L+ I + + AGI+ +G +H E+
 Sbjct: 69 GLATTGKIVPASSPMPASGRAMEIIRNLVAYPKLVKIVATHAGITVGEDGASHQMCD 128
- 40 Query: 126 LGLRLRLPKATIFEVSDDIQFEALKQTLSDGLKLYRTIRKAPVAVYEGRE---DFSK 181
 + ++R IP + +D + +++ G Y+R R+ +YR E + K
 Sbjct: 129 IAIMRAIFNMVVIAPTDYHTQNVIRTLAEYKGFVYVMPRRDTEIYENKEEATPEIGK 188
- 45 Query: 182 GFQILRGKGDITLVAGGIMVSRATEAADYVLEKLGIRASVIDLFKIKPLPEELKPLLDQS 241
 G I L G+D+T++A+G V A+ A +LKE GI A +++ IKP+ EE+ D
 Sbjct: 189 GKI-LVDGEDLTIATGEENVFALRAGEILKENGISAEIVEMATTIKPIDERLIKKSQD-F 246
- Query: 242 IVTIEHNHRIIGGISALCEWL-SNEKDTTVSRMGLIDERPOGVQGMVYLLSEGLAVKDIVQ 301
 +VT+SH+H IGG+G A+ E + S + + R+GI++ FG+ G+ + LL+ YGL + I +
 Sbjct: 247 VVTVEDHSITIGLGRVAEVIASNGLNKKLLRIGINDVPGRSKADELKLYYGDGSEIAK 307

- 50 There is also homology to SEQ ID 520.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 157

A DNA sequence (GBSx0163) was identified in *S. galactiae* <SEQ ID 523> which encodes the amino acid sequence <SEQ ID 524>. Analysis of this protein sequence reveals the following:

55

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Possible site: 24
 >>> Seems to have no N-terminal signal sequence

5 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.2517(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

10 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 158

A DNA sequence (GBSx0164) was identified in *S.agalactiae* <SEQ ID 525> which encodes the amino acid
 15 sequence <SEQ ID 526>. Analysis of this protein sequence reveals the following:

Possible site: 35
 >>> Seems to have no N-terminal signal sequence
 20 INTEGRAL Likelihood = -6.42 Transmembrane 119 - 135 (114 - 145)
 INTEGRAL Likelihood = -5.10 Transmembrane 33 - 49 (32 - 50)
 INTEGRAL Likelihood = -4.30 Transmembrane 94 - 110 (94 - 111)
 INTEGRAL Likelihood = -3.66 Transmembrane 67 - 83 (60 - 83)

 ----- Final Results -----
 25 bacterial membrane --- Certainty=0.3569(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

No corresponding DNA sequence was identified in *S.pyogenes*.

A related GBS gene <SEQ ID 8503> and protein <SEQ ID 8504> were also identified. Analysis of this
 30 protein sequence reveals the following:

Lipop: Possible site: -1 Crend: 4
 SRCFLG: 0
 MOG: Length of UR: 22
 Peak Value of UR: 2.96
 35 Net Charge of CR: 2
 MOG: Discrim Score: 10.55
 GVH: Signal Score (-7.5): -4.31
 Possible site: 22
 >>> Seems to have an uncleavable N-term signal seq
 Amino Acid Composition: calculated from 1
 40 ALOM program count: 6 value: -6.42 threshold: 0.0
 INTEGRAL Likelihood = -6.42 Transmembrane 154 - 170 (149 - 180)
 INTEGRAL Likelihood = -5.10 Transmembrane 68 - 84 (67 - 85)
 INTEGRAL Likelihood = -5.04 Transmembrane 6 - 22 (2 - 24)
 45 INTEGRAL Likelihood = -4.30 Transmembrane 129 - 145 (129 - 146)
 INTEGRAL Likelihood = -3.66 Transmembrane 102 - 118 (95 - 118)
 INTEGRAL Likelihood = -3.56 Transmembrane 29 - 45 (29 - 46)
 PERIPHERAL Likelihood = 0.79 285
 modified ALOM score: 1.78
 50 icml HYPID: 7 CPP: 0.357

 *** Reasoning Step: 3

 ----- Final Results -----
 55 bacterial membrane --- Certainty=0.3569(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

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The protein has homology with the following sequences in the GENPEPT database:

```
>GP:CAE13541 GB:Z99112 ribosomal protein S15 (BS18) [Bacillus subtilis]
Identities = 55/89 (61%), Positives = 71/89 (78%)

5 Query: 1 MAISKEKNEIIAQYARHSGDTGSVEVQVAVLTWEINHLNDHIQHKKHATYRGLMKKI 60
MAI++E+KN++I ++ HE DTGS EVQ+A+LT IN+LN+H++ HKKLN + RGL+K +
Sbjct: 1 MAITQERKNQLINSEFKTHSSDTGSPEVQIAITLDSINHLNEHLTHHKEDHSRGLLNIV 60

10 Query: 61 GHRNLLAVLRRTDVNRYRELIQSLGLRR 89
G RRNLL YLR DV RYRELI LGLRR
Sbjct: 61 GKRRLTYLRNKDVIRYRELIINKLGLRR 89
```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 529> which encodes the amino acid sequence <SEQ ID 530>. Analysis of this protein sequence reveals the following:

```
15 Possible site: 41
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
20 bacterial cytoplasm --- Certainty=0.3746 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

An alignment of the GAS and GBS proteins is shown below:

```
Identities = 88/89 (98%), Positives = 88/89 (98%)

25 Query: 1 MAISKEKNEIIAQYARHSGDTGSVEVQVAVLTWEINHLNDHIQHKKHATYRGLMKKI 60
MAISKEKNEIIAQYARHSGDTGSVEVQVAVLTWEINHLN HIKHKKHATYRGLMKKI
Sbjct: 1 MAISKEKNEIIAQYARHSGDTGSVEVQVAVLTWEINHLNHSIQHKKHATYRGLMKKI 60

30 Query: 61 GHRNLLAVLRRTDVNRYRELIQSLGLRR 89
GHRNLLAVLRRTDVNRYRELIQSLGLRR
Sbjct: 61 GHRNLLAVLRRTDVNRYRELIQSLGLRR 89
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 160

A DNA sequence (GBSx0166) was identified in *S.galactiae* <SEQ ID 531> which encodes the amino acid sequence <SEQ ID 532>. This protein is predicted to be polyribonucleotide nucleotidyltransferase (pnp). Analysis of this protein sequence reveals the following:

```
40 Possible site: 46
>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood = -0.64 Transmembrane 448 - 464 ( 448 - 464)

----- Final Results -----
45 bacterial membrane --- Certainty=0.1256 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
```

A related GBS nucleic acid sequence <SEQ ID 9501> which encodes amino acid sequence <SEQ ID 9502> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```
>GP:AAC43595 GB:U29668 polynucleotide phosphorylase [Bacillus subtilis]
Identities = 428/694 (61%), Positives = 532/694 (75%), Gaps = 4/694 (0%)

55 Query: 7 KQVFEMIFAGKGLVVEVQVAKQANGSVVRYGDSTVLTAAVMSKMSIGDFPLQVNYR 66
```

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K VF + +AG+ L VETQ+AKQANG+V++RYGD+ VL+ A SK+ DFFPL VNYE
 Sbjct: 5 KHVFTIDMAGRTLLIVETQGLAKQANGAVMIRYGDVAVLSTATASKEPKPLDFFPLVNYE 64

Query: 67 EKVYAGKPGPGNPKRGPRSTDATLITARLIDRPIRPMFAGSFRNEVQVINTVLSDEHA 126
 E++YA GK PGGF KREGRPS A L +RLIDRPIR+FA+GFRNEVQV+ V+S D+N
 Sbjct: 65 ERLYAVGKIPGSPFKRGPRSEKAVLASKLIDRPIRPLFADGFRNEVQVISIVMSVDQNC 124

Query: 127 SAPMAAMFOSSLALSISDIPFNGPIAGVAYVVDGHTFIINPTAQDQASALETVAGTK 186
 S+ MAAMFOSSLALS+SDIFF GPIAGV V +D FIINPT + E S + L VAGTK+
 Sbjct: 125 SSEMAAMFOSSLALSVDIPFNGPIAGVTVGRIDQPTINPTVDQLEKSDINLVVAGTKD 184

Query: 187 AINMVEGAKELSEKIMLEALLKGHEAVCELIAPQREIVTAICGKKAVERLLQVLPRL 246
 AINMVE+GA E+ EKIMLEA++ GHE + LIAPQREIV A+GKEK+E++L +D+D EL
 Sbjct: 185 AINMVEAGADEVPKEIMLEAIMPCHSEKIKRLIAPQREIVAAVGKIKSEIKLPEIDELAE 244

Query: 247 EILATHNIALQAQVCEKKAKEAATKAVKEVVGSEYEAHYAEHVEYDRIMRDVAEILBQ 306
 ++ A L A+QV EK ARE A VK V+ ++E EH+E ++ V +IL +
 Sbjct: 245 KVKALAEEDLLAKIQVHEKHAREDAINEVNAVAKFEDE--EHDE--DTIKQVKQLSK 300

Query: 307 MEHAERVLITEKIRPDGRVDEIRPLDAEIDFLPQVHGGSLPTRGQTQALSVLTLAPM 366
 + EVRLITE+K+RPDGR VD+IRPL +E+ LP+ HGGSLPTRGQTQALS V TL +
 Sbjct: 301 LVNNEVRRLITEKVRPDGRGVDQIRPLSSEVGLPRTTRGSGSLPTRGQTQALS VCTLLAL 360

Query: 367 GEAQILIDGLPEYKRPMMHYNFPQSVGEGTRYGAAGRREIGHGALGERALEQVLPRL 426
 G+ QI+DGL E KRPMMHYNFPQ+SVGEGT GRREIGHGALGERALE V+P+
 Sbjct: 361 GNVQILDGLVEESKRPMMHYNFPQSVGEGTGMRCGRREIGHGALGERALEVPIPSEK 420

Query: 427 EFPYAIRLVAEVLSENGSSQASICAGTLAMAGGVPIKAPVAGIANGLSIDGNTVTLT 486
 +FPY +ELV+EVLESNGS+SQASICA TLA+M GVPIKAPVAGIANGL+ G +YTVTLT
 Sbjct: 421 DFPYTVLVLSEVLSENGSTSQASICASTLAMADAGVPIKAPVAGIANGLSGSEHYTLT 480

Query: 487 DIQGLDEHFDMDPKVAGTEGITALQMDIKIGITPQILEEALQAKKARPELLVDLHG 546
 DIQ+ED GMDPKVAGT +G+TALQMDIKIB++ +ILEAL QAKK R EIL+ +
 Sbjct: 481 DIQGNEDALGMDPKVAGTEKGVITALQMDIKIBGLSEILEALQAKKGRMEILMSMLA 540

Query: 547 AIAEPRLQAPTPAKIDIMIKIDVDKIKVIGKGGETIDKILIAETGVKIDDEGNSLPS 606
 ++E R +L+ APKI + I+ DKI+ VIG G+ I+KII ETGVKIDI++G + I S
 Sbjct: 541 TLSESRRLSEKRYAPKILWTINPDKIRDVIGPSCKQINKIIRETOVKIDIEQGTIFIS 600

Query: 607 SDQAIDRTKDIASLVREAKVGEVYHAKVVRIEKFGAFVNLDPKTDALVHISEIAWTRT 666
 +D++ + K II LVRE +VG++Y KV RIEKFGAFV +F D LVHISE+A R
 Sbjct: 601 TDESQNAKAKKIIDELVREVEVQGLYLCKVRIEKGAFVIFSGDKGLVHISELAEV 660

Query: 667 ANVADVLEIGEEDVVKIKIDDKGRVDSKALL 700
 V DV++IG+E+ VKV +ID +GRV+ S KA+L
 Sbjct: 661 GKVEEVVKIGDEILVKVTEIDKQGRVNLSEKAVL 694

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 533> which encodes the amino acid sequence <SEQ ID 534>. Analysis of this protein sequence reveals the following:

Possible site: 28
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.64 Transmembrane 444 - 460 (444 - 460)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.1256 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 631/708 (89%), Positives = 664/708 (93%), Gaps = 2/708 (0%)
 Query: 5 MSKQVFEMIFAGKKLIVVETQGVAKQANGSVVRYGDSTVLTAANVSKEMSTGDFPPLQVN 64
 MSKQ F FAGK LVVE GQVAKQANG+ VVRYGDSTVLTAANVSKEM+TGDFPPLQVN
 Sbjct: 1 MSKQTFPTTFAGKPLVVVEGQVAKQANGATVVRYGDSTVLTAANVSKEMATGDFPPLQVN 60

Query: 65 YEEKVYAGKFGGPMKEGRPSTDRITLTLARLIDRPIRMFAEGFRNEVQVINTVLVSFDE 124
 YEEKVYAGKFGGPMKEGRPSTDRITLTLARLIDRPIRMFAEGFRNEVQVINTVLVS+DE
 Sbjct: 61 YEEKVYAGKFGGPMKEGRPSTDRITLTLARLIDRPIRMFAEGFRNEVQVINTVLVSDE 120

5 Query: 125 NASAPMAAMFGSSSLALSTSDIPFNGPIAGVQVAVVGNPIINPTAGQEGASALELTVAGT 184
 NASAPMAAMFGSSSLALSTSDIPFNGPIAGVQV Y+DG PIIRF ++ EAS LEITVAG+
 Sbjct: 121 NASAPMAAMFGSSSLALSTSDIPFNGPIAGVQVGYIDGFTINPDGQMEASLLELTVAGS 180

10 Query: 185 KEAINDVESGAKELSEDTMLEALLKGHEAVCELIAPQREIVTAIGCEKAVELLQVDEPL 244
 KEAINDVESGAKELSE+IMLEALLKGH+A+ ELIAQGE+IV +GKKKAVELLQVD+L
 Sbjct: 181 KEAINDVESGAKELSEDTMLEALLKGHQAIQELIAPQEQIVAVVGKKEAVELLQVDVDL 240

15 Query: 245 QAEIIATHNIALQAQVVEKKKAREAAATRAVKVVGVEYRARYARHEHYDRIMEDVASEIL 304
 QA+I+A+N LQ AVQVEKKKAREAAATRAVKE+V EYE RYAE E IMEDVASEIL
 Sbjct: 241 QADIVAKYNAQLQKAVQVEKKKAREAAATRAVKVVKAEYEERYAEDENLATHMEDVASEIL 300

20 Query: 305 EQMNEHAEVRLITEDIKIRPDGRRVDEIRPLDAEIDPLPQVHSGSLFTRGOTQALSVLTLA 364
 EQMNEHAEVRLITEDIKIRPDGR++DEIRPLDA +DPLP+VEGSGSLFTRGOTQALSVLTLA
 Sbjct: 301 EQMNEHAEVRLITEDIKIRPDGRKIDEIRPLDAVDPLPQVHSGSLFTRGOTQALSVLTLA 360

25 Query: 365 PMGEAQIIDGLPEYKKRPMHYNFPQYSGVGETGRYGAAGREIGHGALGERALEQVLP 424
 PMGE QIIDGL PEYKKR+HYNFPQYSGVGETGRYGAAGREIGHGALGERALEQVLP
 Sbjct: 361 PMGETQIIDGLAPEYKKRFLHYNFPQYSGVGETGRYGAAGREIGHGALGERALEQVLP 420

30 Query: 425 LEEFPYAIRLVAEVLSENGSSQASICAGTALMAGGVPIKAPVAGIAMGLISDGTNYTV 484
 LEEFPYAIRLVAEVLSENGSSQASICAGTALMAGGVPIKAPVAGIAMGLISDGTNYTV
 Sbjct: 421 LEEFPYAIRLVAEVLSENGSSQASICAGTALMAGGVPIKAPVAGIAMGLISDGTNYTV 480

35 Query: 485 LDTIQGLEHFGMDMPKVACTREGITALQMDIKIGITPQILEEALQAKKARFELDLVL 544
 LDTIQGLEHFGMDMPKVACTREGITALQMDIKI GITPQILEEALQAKKARFELDLV+
 Sbjct: 481 LDTIQGLEHFGMDMPKVACTREGITALQMDIKIGITPQILEEALQAKKARFELDLVI 540

40 Query: 545 HGAIAEPFRQLAPAPKIDMKIIVDKIKVVGKGGETIDKIIAETGVKIDIDDEBNVSI 604
 IAEPR+LAPAPKID IKIDVOKIIVVGKGGETIDKIIAETGVKIDID+EGNVSI
 Sbjct: 541 EATIAEPFRQLAPAPKIDTIIKIDVOKIIVVGKGGETIDKIIAETGVKIDIDDEBNVSI 600

45 Query: 605 FSSDQAIDRTKDIASLVREAKVGEVYHAKVVRIEKFQAFVNLFDKTDALVHISIAWT 664
 +SSDQAIDRTK+IIA LVREAKVGEVYHAKVVRIEKFQAFVNLFDKTDALVHISIAWT
 Sbjct: 601 YSSDQAIDRTKEIIAGLVREAKVGEVYHAKVVRIEKFQAFVNLFDKTDALVHISIAWT 660

Query: 665 RTANVADVLEIGEEVDVVKIKIDDKGRVDASMKALLPRPKADNPKKE 712
 RT NV+DVLE+GE+VDVVKIKID+KGRVDASMKALL+PRPK + KKE
 Sbjct: 661 RTTNVSDVLEIGEEVDVVKIKIDEKGRVDASMKALLPRPKPE--KKE 706

45 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 161

A DNA sequence (GBSx0167) was identified in *S.agalactiae* <SEQ ID 535> which encodes the amino acid sequence <SEQ ID 536>. Analysis of this protein sequence reveals the following:

50 Possible site: 39
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1293 (Affirmative) < succ>
 55 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 537> which encodes the amino acid sequence <SEQ ID 538>. Analysis of this protein sequence reveals the following:

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Possible site: 38

>>> Seems to have no N-terminal signal sequence

INTEGRAL Likelihood = -0.43 Transmembrane 83 - 99 (83 - 99)

----- Final Results -----

bacterial membrane --- Certainty=0.1171(Affirmative) < succ>

bacterial outside --- Certainty=0.0000(Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

An alignment of the GAS and GBS proteins is shown below:

Identities = 172/248 (69%), Positives = 211/248 (84%)

Query: 1 MTSTNKLDIRLRAFINAPDNFLDSIGLVNALHSHSTVWASKRFPYAIQVQGVVVFVDTIT 60
 MT +NELDIRLRAFINAPDNFLDS+ LVNA H+ VWA+KEFY I+V+G +V FVFTD
 Sbjct: 1 MTKSNELDIRLRAFINAPDNFLDSLALVNAFHNFPVWAARFPYVIEBVGKVTFTVDKE 60

Query: 61 DLNHFKEBQESARDMFWSRRSLDVLDRASHGLAGLVYNNLKKKGDPGNSTIFYCEDMVQ 120
 D+ FKEBQ+SA+ +W R +L VL+E I+ G AGL++NLKK+GDPGNSTIF DN+Q
 Sbjct: 61 DMARFKEBQKSAQSQYWLERSALAVLEEVITSGAAGLIFNLKKKGDPGNSTIFKSSDMIQ 120

Query: 121 FMNNYTTIILNQLLNEDNIVADIMDKTYLVPAPVHPREBGSFDRLFPTMTPEGKSYVVF 180
 FMN+YTT+LN L++DN+ AD M+K YLVPAPV+P++ +DRLFPTMTPEGKSYVP F
 Sbjct: 121 FMHNYTTVILNQLSDDNVAADTMEKVTYLPAPVYPKDMNHYDLRFP:MTPEGKSYVPAF 180

Query: 181 SNLLSFEKWNHNDPFGAFRKAQGVILAWTIDDIYKPRNGENEIDDTFGVAINPDEQVQ 240
 SNL SF KWYN +DPGG FRKA+GVIL WTIDDIY+PRNGENE+D+TFGVAINPFD+QQ+
 Sbjct: 181 SNLQSFARKWYNQDQDGGFLRKAEGVILWTIDDIYQPRNGENELDETFGVAINPFDQCI 240

Query: 241 LVDWSQVE 248
 LVDWS+++
 Sbjct: 241 LVDWSELD 248

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 162

A DNA sequence (GBSx0168) was identified in *S.agalactiae* <SEQ ID 539> which encodes the amino acid sequence <SEQ ID 540>. This protein is predicted to be serine acetyltransferase (cysE). Analysis of this protein sequence reveals the following:

Possible site: 39

>>> Seems to have no N-terminal signal sequence

INTEGRAL Likelihood = -2.02 Transmembrane 150 - 166 (147 - 168)

----- Final Results -----

bacterial membrane --- Certainty=0.1808(Affirmative) < succ>

bacterial outside --- Certainty=0.0000(Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 9503> which encodes amino acid sequence <SEQ ID 9504> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:CA571304 GB:A130879 serine acetyltransferase [Clostridium
 sticklandii]

Identities = 92/169 (54%), Positives = 125/169 (73%)

Query: 9 KESIAIVKEQDPAARSSLEVLITYPGI:KALAAHRLSHFLWNHNFKLLARMHSQFWRFTWQ 68

-240-

KE+I + +E+DPAA+ ++ +++ PGI A+ HR++H L+N +AR+ SO RF T
 Sbjct: 20 KETISVAREKDPAAKGAINLIVNTPGIHALMFHRVARSLYNRKHFFIARLISQISRFITG 79

Query: 69 IEIHFGATISRGVFDHGSGLVIGETAIVEKGAMLYHGVTLGSTGKDKGERHPTIRKGA 128
 5 IEIHFGA I FIDHG G+VIGETA + ML+H VTLGGTGKDKGERHPT+ +
 Sbjct: 80 IEIHFGAIGRRFIDHGMGVVIGETASIGEDVMLFQVTLGGTGKDKGERHPTVENNVI 139

Query: 129 ISASHSQITGPIEVGENAKVGAAAVVLADVPADVTVVGVPKVVVRHQQ 177
 10 ISA ++GPI +GEN+K+GA AVVL D+P + T VG+PAKVVR+G+K
 Sbjct: 140 ISAGVKVIGPITVIGENSKIGANAVVLHDIPKNATAVGIPAKVVRINGEK 188

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 541> which encodes the amino acid sequence <SEQ ID 542>. Analysis of this protein sequence reveals the following:

Possible site: 35
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.0141 (Affirmative) < succ>
 20 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 162/193 (83%), Positives = 176/193 (91%)

Query: 5 MGNWKESTIAIVKEQDPAARSSIEVILTYPGIKALAAHRLSHFLWNNPKLLARMHQSQWR 64
 25 MGNWKESTIAIVK DPAAR+SIEVILTYPGIKALAAHRLSHFLW H+FKLLARMHQSQWR
 Sbjct: 1 MGNWKESTIAIVKADPAARNSIEVILTYPGIKALAAHRLSHFLWRHHPKLLARMHQSQWR 60

Query: 65 FWTQIEIHFGATISRGVFDHGSGLVIGETAIVEKGAMLYHGVTLGSTGKDKGERHPTIR 124
 30 FWTQIEIHFGA I+ GVFDHG+GLVIGETAIVEKG MLYHGVTLGSTGKDK GERHPT+R
 Sbjct: 61 FWTQIEIHFGAIGLVFDHGSGLVIGETAIVEKGAMLYHGVTLGSTGKDKGERHPTVR 120

Query: 125 KQALISASHSQITGPIEVGENAKVGAAAVVLADVPADVTVVGVPKVVVRHQQKDDLSQRS 184
 35 +GALISAH+Q+IGPI++G NAKVGAAAVVL+DVP DVTVGVPK+VVRHQQK+ QI+S
 Sbjct: 121 QGALISAHQVIGPIDIGANAKVGAAAVVLSDVPEDVTVGVPKVVVRHQQKNNRQIQS 180

Query: 185 IEHIDREESYYSK 197
 ++ RE SY SK
 40 Sbjct: 181 LQKQREVSYQLSK 193

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 163

A DNA sequence (GBSx0169) was identified in *S.agalactiae* <SEQ ID 543> which encodes the amino acid sequence <SEQ ID 544>. Analysis of this protein sequence reveals the following:

Possible site: 29
 >>> May be a lipoprotein
 INTEGRAL Likelihood = -5.89 Transmembrane 32 - 48 (29 - 49)

----- Final Results -----
 bacterial membrane --- Certainty=0.3357 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 164

A DNA sequence (GBSx0170) was identified in *S. galactiae* <SEQ ID 545> which encodes the amino acid sequence <SEQ ID 546>. This protein is predicted to be cysteinyl-tRNA synthetase (cysS). Analysis of this protein sequence reveals the following:

Possible site: 46
>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.2227 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP: CAB11870 GB: Z99104 cysteinyl-tRNA synthetase [Bacillus subtilis]
Identities = 246/465 (52%), Positives = 322/465 (68%), Gaps = 23/465 (4%)

Query: 2 IKIYDVTMRSLQDFIPLEBKGVMVVOGPTVYNYIHIGNARSVVAFOTIRRYFYGQYQV 61
I +Y+T+TR + F+FL BGV MYVOGPTVYNYIHIGNAR +DT+R Y BY GY V
Sbjct: 3 ITLNYTLTRQKETPFLBEGKVVMVVOGPTVYNYIHIGNARPAIVDTVENYLEYGYDV 62

Query: 62 NYISNFTOVDDKIIKGAARAGMOTKFSFDFISAPMEDVAALGVKPKATNPRVIDYMDI 121
Y+SNFTOVDDK+IK A E G D + S++FI A+ BUV ALG + A +PRV++ MD I
Sbjct: 63 QVSNFTOVDDKLIKAAELGEDVPTISRFPIKAYFEDVGALGCRKADLHPRVNMDAI 122

Query: 122 IDPVKVLVDKEFAYRANGDYVFRVSKSHHYAKLANKTLEDLEIGASGRVDGSGIKENFL 181
I+FV LV K +AYE+ GDVYF+ Y KL+ ++++L GA RV GR KE+ L
Sbjct: 123 IDFVDQVWKKGYAYESBGUVTYFKTRAFEGYGLKSGSIDELRSGARIV---GEKKDAL 179

Query: 182 DFLMWKSAKSGEVSWSWSPWGKGRPGWHIECSVMATEILGUTIDIHGGGADLEFPHTNEI 241
DFALMW+AK GE+SW+SPWGKGRPGWHIECS M + LGD IDIH GG DL FPH NEI
Sbjct: 180 DFLMWKAAKEGELISWDSFWGKGRPGWHIECSAMVKKVLGDQIDIHAGQDLTFPHNEI 239

Query: 242 AQSEAKTGKTFANYWHNGFPVNDNEKMSKSLGNFTVHDMLKSVSGQVIRFPLATQQYR 301
AQSEA TGKTF YW+HNG++N+DNEKMSKSLGNF+ VHD++K D Q++RFF+ + YR
Sbjct: 240 AQSEALTGKTFKAYWIRNGYINIDNEKMSKSLGNFVLVHDIKQHPQLLRPFMLVHYR 299

Query: 302 KPVNPTKAVHDABEVNLKYLKNTF-----NLPIQENANDEELEGFKVAFQGM 350
P+N++E+ + + + LK + NL ++ E++E+ KAF+ MD
Sbjct: 300 HPINYSBELLNTEKASPRLLKATYASNLQRLNSNTLTDQQMLKVEHEHKAPEEMD 359

Query: 351 DDFNTANGITVIFEMAKWIN-----SGHYTSRVKETFAELLEIFGI-VFQEEVLDA 401
DDFNAN I+V+P++AK N + H + E F ++ + G ++E+LD +
Sbjct: 360 DDFNTANISVLFLAKHANYTLQKDHDTADHTVITAFIEMPDRIVSGLFSLGBOELLQ 419

Query: 402 IESLIEQRQEARANRDFATADRINDELAKQGIKLLDTKDGVRWR 446
IE LIE+R EAR NEDFA +D+IRD+L I L DT G RW R
Sbjct: 420 IEDLIEKRNKARNRDFALSIDIRDLQKSMNIILEDTAQGRWKR 464

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 547> which encodes the amino acid sequence <SEQ ID 548>. Analysis of this protein sequence reveals the following:

Possible site: 46
>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.1765 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 357/447 (79%), Positives = 401/447 (88%)

```

5  Query: 1 MIKIYDITMTRSLQDFIPLNKGKVMYVCGFTVYNYIHIGNARSVAVFDITIRRYPEYGYQ 60
    MIKIYDITMTRSL+P+PL E VN+YVCGFTVYNYIHIGNARS VAFDITIRRYPEY GYQ
    Sbjct: 1 MIKIYDITMTRSLRKFVPLTEFTVNIYVCGFTVYNYIHIGNARSVAVFDITIRRYPEYGYQ 60

    Query: 61 VNVISNFTDVDDKIIKGAAGMDTKSPDKFISAPMEDVAALGVKPKATKNRPVIDYMD 120
    VNVISNFTDVDDKIIK A +AG+ K SD+FI+AF+RD AIGVKPAT+NDPV+DY+ E
10  Sbjct: 61 VNVISNFTDVDDKIIKATQAGVSPKELSDRFIAAFIEDTKALGVKPKATNDPV+MOYIAR 120

    Query: 121 IIDPVKVLVDKGFAYEANGDVYPRVSKSHHYAKLANKITLEDLEIGASGRVDGGEIKENP 180
    II FV+ L++K+PAYRA+GDVYPRV KS HYAKLANKITL+LE+GASGR D E +KENP
    Sbjct: 121 IISFVBSLIEKDPAEADGDVYPRVSKSHHYAKLANKITL+SELEVGASGRD+ETALKENP 180

15  Query: 181 LDFALWKSAGSGEVSWSRSPWGKRGPGWHIECSVMATEILGDTIDHGGADLEFPFHITNE 240
    LDFALWKSAG+GEVSW+SPWG GRPGWHIECSVMATEILGDTIDHGGADLEFPFHITNE
    Sbjct: 181 LDFALWKSAGGEVSWSDSPWGKRGPGWHIECSVMATEILGDTIDHGGADLEFPFHITNE 240

    Query: 241 IAQSEAKTGKTFANYWMHNGFVYVNEHMSKSLGNFITVHMLKSVDDGVIRFPLATQQY 300
    IAQSEAKTGKTFANYWMHNGFV VNEHMSKSLGNF+TVHML++VDQGV+RFPLATQQY
    Sbjct: 241 IAQSEAKTGKTFANYWMHNGFVYVNEHMSKSLGNFITVHMLQTVDDGVIRFPLATQQY 300

    Query: 301 RKP+NFTEKAVHDAEVLNLYLKNTFNLPIQNANDEELQFVKAFQGMDDDFNTANGIT 360
    RKP+NFTEK +HDAE+NLKYLKNT P+ E A+++EL+QFV AFO AMDDDFNTANGIT
    Sbjct: 301 RKP+NFTEKTHDAEVLNLYLKNTLQQLTFETADQELQFVIAPQGMDDDFNTANGIT 360

    Query: 361 VIFEMAKWINGSGHYTSRVKTFASLLKIPGVQFEVLOADIESLIFQEQEARNRDPAT 420
    V+FAKAWINGS YI VK F ++L +PGI+P+EEVL+ DIK+LI +RQEARNRDPAT
    Sbjct: 361 VVDFMAKWINGSYITEPVKSAFEIMLAVFGIIFEEVLEVDIEALIAKRGQEARNRDPAT 420

    Query: 421 ADRIREDLAKQGIKLLDTKDGVRWTRD 447
    AD IRD+LA QGIKLLDTKDGVRW RD
    Sbjct: 421 ADAIRDQLAVQGIKLLDTKDGVRWLRD 447
35

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 165

A DNA sequence (GBSx0171) was identified in *S. agalactiae* <SEQ ID 549> which encodes the amino acid sequence <SEQ ID 550>. Analysis of this protein sequence reveals the following:

```

    Possible site: 53
    >>> Seems to have no N-terminal signal sequence

    ----- Final Results -----
45  bacterial cytoplasm --- Certainty=0.0259 (Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
    bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 9505> which encodes amino acid sequence <SEQ ID 9506> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:CA11871 GB:299104 similar to hypothetical proteins [Bacillus subtilis]
Identities = 58/122 (47%), Positives = 87/122 (70%)

55  Query: 3 DVRLINGIALAFGIDAVVSLYIRRHILMQGFTKPNQLHRKATQYVSANQAALLINAMLE 62
    D + +NG+ALA+ GDA++ +Y+R HL+ QGFTKPN LH+K++ VSA +QA ++ + +
    Sbjct: 9 DSKQKLALALAYIGDAIPEVYVRHLLKQGFTKPNLHKSSRIVSAKSAQRIFLPLQ 68

    Query: 63 NILTDEBQLIYKKGKRNASHITKAKNADITTYRMSYSGPEALMG+LDMTGQIKRIEFTLQWC 122

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+ T+E+ + KRGRNA S T KN D+ TYR ST FEAL+GYL + + +RL L+
 Sbjct: 69 SFTTEEEAVLKRGRNAKSGTTPKNTDVTQITRYSTAFEALLGYLFLKKKEERLSQAVAE 128
 Query: 123 IE 124
 ++
 Sbjct: 129 IQ 130

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 551> which encodes the amino acid sequence <SEQ ID 552>. Analysis of this protein sequence reveals the following:

Possible site: 56
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 99/127 (77%), Positives = 111/127 (86%)
 Query: 2 IDVRLINGIALAFEGSDAVSYLYIRPHLMQGFYKPNQLHRKATQVSNAAQALLINAMLE 61
 +DV LINGIALAFEGSDAVYS Y+RPHLI Q3 TKP+QLHR AT+YVSA AQA LI AMLE
 Sbjct: 5 VDVLNGLIALAFEGDAVYSYVVRHLI PQGKTKPSQLHRLATRYVSAQAQANLIQAMLE 64
 Query: 62 ENILATDEQLIYKGRNANSHTKAKNADIITYRMSTGFALMGYLDWGTQIKRLETLIQW 121
 +LT++E+ IYKGRN NSHTKAKNADIITYRMSTGFER+MGYLDM GQ +RLE LI+W
 Sbjct: 65 AQLLEKSEDIYKGRNTNSHTKAKNADIITYRMSTGFALMGYLDWGTQIKRLETLIQW 124
 Query: 122 CIEITEK 128
 CIE +EK
 Sbjct: 125 CIEIVYK 131

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

35 Example 166

A DNA sequence (GBSx0172) was identified in *S.galactiae* <SEQ ID 553> which encodes the amino acid sequence <SEQ ID 554>. This protein is predicted to be spoU rRNA methylase family protein. Analysis of this protein sequence reveals the following:

Possible site: 30
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1478 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAB11872 GB:Z99104 similar to hypothetical proteins [Bacillus subtilis]
 Identities = 113/244 (46%), Positives = 163/244 (66%), Gaps = 6/244 (2%)
 Query: 11 ESSDLVYGLHAVTFESLRANIG-NKLYLQDDLKGNVVKVICALATEKKVVISMTPKKTLSD 69
 + D V G +AV E+L+++ KL++ ++ +V LA ++ ++I +P+K L
 Sbjct: 3 QQSDYVIGKNVAVITLTKSDRKLYKLMAENIVKQQAQVIELAKKQGTITQVPRKLDQ 62
 Query: 70 MINGGVHGGFVLKVFAYADLSHIMTKAENE-ENPLILILDGLTDPHNLGSLRTADAT 128
 M G HGV V +V+ + YA+L ++ AR + E P LIID L DPHNLGSI+RTADA
 Sbjct: 63 NVTOG-QHGVVAQVAAYEYAEIDDLVYKAAEKKNEQPPFLLIDLELDPHNLGSI+RTADAV 121

-244-

Query: 129 NVTGIIIPKHSVGVTPVVSKTSTGAVEHVPIARVTNLSQTLTLDKKEFWIPGTMNGT 188
 GI+IPK R+VG+T V+K STGA+EH+P+ARVTNL+TL+ +K++ W+ GTD +
 Sbjct: 122 GARGIIVIPKRSVGLTITVAKASTAIEHIPVARVTNLARTLEEMKRGIVVGTADASR 181

5 Query: 189 PSKHWNTKGK--LALVIGNEGKISHNIKKQVDEMIIIPNGHVQSLNASVAARAILMYEV 246
 + N G LALVIG+BGKG+ +K++ D +I +PM G V SINAGVAA +LMYEV
 Sbjct: 182 EDPR-NMDXNMLPALVIGSBGKGMRLVREKCDFLIKLPMGKVTSLNASVAAGILMYEV 240

10 Query: 247 FRNR 250
 +R R
 Sbjct: 241 YRER 244

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 555> which encodes the amino acid sequence <SEQ ID 556>. Analysis of this protein sequence reveals the following:

15 Possible site: 36
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1037 (Affirmative) < succ>
 20 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 206/248 (83%), Positives = 225/248 (90%), Gaps = 1/248 (0%)

25 Query: 3 NKDKQPKRESSDLVYGLHAVTESLRANTGNKLYLQDLRGKIVDKVALATEKKVSIWT 62
 N+DK E++D+VYG+HAVTESL+ANTGNKLY+Q+DLRGK VD +K+LAT+KKV+ISWT
 Sbjct: 10 NEDKD-TIETNDIVYGVHAVTESLQANTGNKLYIQDLRGKVDNKLATQKQVAISWT 68

30 Query: 63 PKKTLSDMTNGVGHQGFVLKVSERFAYADLSRMTKAENEEMPLLILLDGLTDPHNLGSIL 122
 PKKTLs MT+G VHQGFVL+VS FAY D+ EI+ AR E NPLILLDGLTDPHNLGSIL
 Sbjct: 69 PKKTLQMTDGAHVQGFVLKVSFAFYTDVDEILEIABQANPLLILLDGLTDPHNLGSIL 128

35 Query: 123 RTADATNVGIIIPKHSVGVTPVVSKTSTGAVEHVPIARVTNLSQTLTLDKKEFWIPG 182
 RTADATNV G+IIPKHSVGVTPVVSKTSTGAVEH+PIARVTNLSQTLTLDKKEFWIPG
 Sbjct: 129 RTADATNVGVIIIPKHSVGVTPVVSKTSTGAVEHPIARVTNLSQTLTLDKKEFWIPG 188

Query: 183 TDMNGTTPSKHWNTKGK--LALVIGNEGKISHNIKKQVDEMIIIPNGHVQSLNASVAARAIL 242
 TDMNGTTPS WNT GKALVIGNEGKIS NIKKQVDEMIIIPNGHVQSLNASVAARAIL
 40 Sbjct: 189 TDMNGTTPSDWNTNGKALVIGNEGKISNIKKQVDEMIIIPNGHVQSLNASVAARAIL 248

Query: 243 MYEVFRNR 250
 MYEVFRNR
 Sbjct: 249 MYEVFRNR 256

45

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 167

A DNA sequence (GBSx0173) was identified in *S.agalactiae* <SEQ ID 557> which encodes the amino acid sequence <SEQ ID 558>. Analysis of this protein sequence reveals the following:

55 Possible site: 18
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.2187 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

-245-

>GP:CAB11873 GB:Z99104 similar to hypothetical proteins [Bacillus subtilis]
Identities = 67/147 (45%), Positives = 94/147 (63%), Gaps = 2/147 (1%)

Query: 6 ILLVDGYNMIAFWKDTQLPKFNRLAEAREVLLRLKLNHYAHFEHIDICVFDQYVGVVR 65
ILLVDGYNMI W + L K-N EEAR+VL+K+ Y + +I VFDA V G+
Sbjct: 3 ILLVDGYNMIGAWPQLKDL-KANSFEAREVLIQMAQYQSYTKNRVIVVFDHLVKGLE 61

Query: 66 QRYDQYKISVIFTEEDTADSYIERAAALNQSVLNLVSATSDIANEQWTFISQCALRVS 125
++ +++ VIPT+E+ETAD IK+ A LN ++ + VATSD EQW IF QCALR S
Sbjct: 62 KKQTNHREVIPTKENTADERIEKLAQALN-NIATQIHVATSDYTEQNAIQCALRKS 120

Query: 126 ARELEQRVATVKSDDLDMSSQIDLSTP 152
AREL + V T++ +++ +I P
Sbjct: 121 ARELLREVETIERRIERRVRKITSEK 147

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 559> which encodes the amino acid sequence <SEQ ID 560>. Analysis of this protein sequence reveals the following:

Possible site: 46
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.2465 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 130/167 (77%), Positives = 149/167 (88%), Gaps = 1/167 (0%)

Query: 3 KHSILLVDGYNMIAFWKDTQLPKFNRLAEAREVLLRLKLNHYAHFEHIDICVFDQYVP 62
K ILLVDGYNMIAFW+ TROLFK+N+L+AR LL KLNHYAHFE+I+IICVFDQYVP
Sbjct: 2 KKRILLVDGYNMIAFWQSTRLQFKTNQLDQARNVLLTYLKNHYAHFENITICVFDQYVP 61

Query: 63 GVRQRYDQYKISVIFTEEDTADSYIERAAALNQSVLNLVSATSDIANEQWTFISQCAL 122
G+RQRYDQY IGW+PTEEDTADSYIER AELN + +++V VATSDIANEQWTFISQCAL
Sbjct: 62 GLRQRYDQYIISVVPTEEDTADSYIERAAELN-TAHMVEVATSDIANEQWTFISQCAL 120

Query: 123 RVSAARELEQRVATVKSDDLDMSSQIDLSTPFLKRPANDEQLKLDPL 169
RV+ARELEQRV TVK+DLDMSS IDL TPKLR++ QL +LKDF+
Sbjct: 121 RVTARELEQRVHTVKDDLDMSSRIDLKTFLKRPFDQQLQLKDFM 167

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 168

A DNA sequence (GBSx0174) was identified in *S.galactiae* <SEQ ID 561> which encodes the amino acid sequence <SEQ ID 562>. Analysis of this protein sequence reveals the following:

Possible site: 58
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.4889 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAB12951 GB:Z99109 yits [Bacillus subtilis]
Identities = 100/284 (35%), Positives = 157/284 (55%), Gaps = 6/284 (2%)

Query: 1 MTPKILIDTSSTEDLEKWAQEHNVDIIGLTIRLDGKTYRTVDEKITSDFILERMQEGALP 60
MT ++ DS +DL + +E + I L + L K +E I +D + E MQ G P

-246-

Sbjct: 1 NTVHLIADSATDLPRSPYPERKGIGFPLRVSLGDKPEFDA--VTIHADQIPEAMQGETP 58
 Query: 61 TTSQINVGQFEFVSTYAENDHALLYLALSSHLGTYQSATAREMVLDKYDPAQIEIVD 120
 TSQ + + VF YAE LY-A SS LSGTYQ-A + V +++PD + +VD
 Sbjct: 59 KTSQASPTQIKRVPLQYAFETGDPALYIAPSSGLSGTYQTAVMLANSVGEFPDOLRVID 118
 Query: 121 TAAASGEGVLAMLAATKREQSKSLREVVKQKIESLLPKLNTYFLVDDLNLHMRSGRLSKG 180
 + AS G G+ A G +++E+ + + + +L F VDDL +L R GR+SK
 Sbjct: 119 SKCASLGYGLAVRHAADLCINMENTIQEITSVGNPQSLEHIFTVDLTYLARGGRISKT 178
 Query: 181 AALIGSVAKIKLLKLDSEGLVFPKTRGRKKGLK--EIVTQATKLSYSTLIAYSG 237
 +A +G + IKELL++ +GKLVP K RG+K K E++ + S T+ I+Y+
 Sbjct: 179 SAFPVGILNIKPLQME-DGKLVPLEKIRGQKLPKRIELMKERGDMSQYTVGISYAA 237
 Query: 238 EKDSAQVMEQLLADERIEKVIIRPLGPVISAHVSGSALALPSL 281
 K- A MK + + +E+I+ P+ I +H G G IAF L
 Sbjct: 238 NKEKATYMKHILIEAPFKKEIIMHPISSAIGSHAGPOTLAIFFL 281

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 563> which encodes the amino acid sequence <SEQ ID 564>. Analysis of this protein sequence reveals the following:

Possible site: 18
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3247(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 167/286 (58%), Positives = 227/286 (79%)
 Query: 1 NTFKILIDSTSDLDHQAQENVDILGILTELDGTYETVGEDEKITSDFILLERMORGAKP 60
 NTF I+TDST+DL++ WA++H++ +IGLTI DG+ YETVG +I+SD+LL++M+ G+ P
 Sbjct: 1 NTFITIMDSTADLNQTAESDHDIVLIGLITLDCGEVYETVGNRISSDYLLKKGKAGSHP 60
 Query: 61 TTSQINVGQFEFVSTYAENDHALLYLALSSHLGTYQSATAREMVLDKYDPAQIEIVD 120
 TSQINVG+FE+VF +A N+ ALLYLA SS LSGTYQSA +AR++V + YPDA IRIVD
 Sbjct: 61 QTSQINVGFEKVFREHARNKALLYLAPSSVLSGTYQSALMARDLVREYDPAVIEIVD 120
 Query: 121 TAAASGEGVLAMLAATKREQSKSLREVVKQKIESLLPKLNTYFLVDDLNLHMRSGRLSKG 180
 T+AA+ GEG L +LA + R GK+L E K +E++++P+L TYPLVDDL HLMR GRISKG
 Sbjct: 121 TLAAGGGEYVLTILAAEARDSGNLLTETKDIVEAVIPRLRTYPLVDDLPHMRGGRGLSKG 180
 Query: 181 AALIGSVAKIKLLKLDSEGLVFPKTRGRKKGLKIEIVTQATKLSYSTLIAYSGKD 240
 +A +GS+A IKELL +D EGKLVF AK RGR+K IKE+V Q K ++ ST+I++Y+ ++
 Sbjct: 181 SAFPGLASIKPLMWDERGKLVPKIKRGRKAIKEMVAQVEKDIASTVIVSVTSDQG 240
 Query: 241 SAQVMEQLLADERIEEVIIRPLGPVISAHVSGSALALPSLGEENR 286
 SA+ ++E+LA E I +V++ PLGPVISAHV LA+P +G+ +R
 Sbjct: 241 SAEKLRZELIAHENISDVMPLGPVISAHVGVNTLAVFVIGQNSR 286

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 169

A DNA sequence (GBSx0175) was identified in *S.agalactiae* <SEQ ID 565> which encodes the amino acid sequence <SEQ ID 566>. Analysis of this protein sequence reveals the following:

Possible site: 56
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -8.76 Transmembrane 43 - 59 (40 - 62)

-247-

----- Final Results -----

bacterial membrane --- Certainty=0.4503(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

5

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

10 **Example 170**

A DNA sequence (GBSx0176) was identified in *S.galactiae* <SEQ ID 567> which encodes the amino acid sequence <SEQ ID 568>. This protein is predicted to be ribosomal protein L13 (rplM). Analysis of this protein sequence reveals the following:

Possible site: 55

15

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3426(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

20

A related GBS nucleic acid sequence <SEQ ID 9507> which encodes amino acid sequence <SEQ ID 9508> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

25

>GP:BAB03887 GB:AP001507 ribosomal protein L13 [Bacillus halodurans]
 Identities = 89/144 (61%), Positives = 113/144 (77%)

Query: 36 KTTFMKPGQVERKWTYVDAADVLGRLSAVVASVLRGHNKPTTTHDITGDGVIVINAE 95
 +TT+MAKP +VERKWTYVDA LGRLL+ VAS+LRGK+KPT+TTH DTGD VI+INAE

30

Sbjct: 2 RTTTFMKPNEVERKWTYVDAEQTGLRLASEVASILRGHKKFTYTHVDTGDHVIINAE 61

Query: 96 KVLTGKKASDKIYYTHSMYPOGLKQISAGELSKNVRLEKSVKMLPHHTLGRAGQM 155
 K+ LTK K DKIIY HS +POGLK- A ++R+ +++E ++KMLP NTLGR QGM

35

Sbjct: 62 KIHLTGNKLQKIKIYYRHSCHPGGLKETRAADMRANKPRIMLELAIKMLPKHTLGRKQGM 121

Query: 156 KLVVFGGEHTHAQQPEVLDSG 179

KL V+ G EH H AQ+PEV ++ G

Sbjct: 122 KLVVYAGSEKHQAQKPEVYELRG 145

40

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 569> which encodes the amino acid sequence <SEQ ID 570>. Analysis of this protein sequence reveals the following:

Possible site: 57

>>> Seems to have no N-terminal signal sequence

45

----- Final Results -----

bacterial cytoplasm --- Certainty=0.4249(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

50

An alignment of the GAS and GBS proteins is shown below:

Identities = 167/184 (90%), Positives = 171/184 (92%), Gaps = 4/184 (2%)

Query: 1 NPTFPVVRPNLNLWTLVDNRNHT--CKQ-KRIRIGRIMNKTTFMKPGQVERKWTYVDAAD 57
 +TPTFP RPNL NT D H CKQ RIRIGRIMNKTTFMKPGQVERKWTYVDAAD

-248-

Sbjct: 1 LPTTPFRPRNLPTT7-DGTEHPSPCKQLIRIGEIMNKITTFMAKPOQVERKWVVDAD 59
 Query: 58 VPLGRLSAVVASVLRGKKNKPTTPTHTDTGDFVIVDAEKVLTGKKASDKIYYTHSNYPG 117
 VPLGRLEAVVASVLRGKKNKPTTPTHTDTGDFVIVDAEKVLTGKKA+DK+YYTHSNYPG
 Sbjct: 60 VPLGRLSAVVASVLRGKKNKPTTPTHTDTGDFVIVDAEKVLTGKKATDKVYYTHSNYPG 119
 Query: 118 GLKQIQRGELRSKNVRLIEKSVKGMLEPHNTLGRAQGMKLVFVUGBHTHAQAQPEVLDI 177
 GLK I+AGELRSKNVRLIEKSVKGMLEPHNTLGRAQGMKLVFVUGBHTHAQAQPEVLDI
 Sbjct: 120 GLKSTAGELRSKNVRLIEKSVKGMLEPHNTLGRAQGMKLVFVUGBHTHAQAQPEVLDI 179
 Query: 178 SGLI 181
 SGLI
 Sbjct: 180 SGLI 183

- 15 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 171

A DNA sequence (GBSx0177) was identified in *S. agalactiae* <SEQ ID 571> which encodes the amino acid sequence <SEQ ID 572>. This protein is predicted to be 30S ribosomal protein S9 (rpsI). Analysis of this protein sequence reveals the following:

Possible site: 53
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1761 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAB11926 GB:Z99104 ribosomal protein S9 [Bacillus subtilis]
 Identities = 88/130 (67%), Positives = 105/130 (80%)
 Query: 1 MAQAQYAGTGRRKNVAVRVLVPGTGKITINKDVEEYIPHALLELVINQPFVAVTSTQGS 60
 MAQ QY GTGRRK++VARVELVPG G+I +N +++ E+IP A L I QP +T T G+
 Sbjct: 1 MAQVQYYTGTRRRKSSVAVRVLVPGEGRIVVNNREISEHIPSAALEDIKQPLTLTETAGT 60
 Query: 61 YDVFVNVGGSYGQSGAIRHGIRALLEVDPDFRDLKRAGLLTRDARMVERKKPGLKX 120
 YDV VNV GGS +GQ+GAIRHGI+RALLE DP++R +LKRAGLLTRDARM ERKK GLK
 Sbjct: 61 YDVLNVHGGGLSGQGAIRHGIRALLEADPEYRTTLKRAGLLTRDARMERKKYGLKG 120
 Query: 121 ARKASQFSKR 130
 AR+A QFSKR
 Sbjct: 121 ARRAQFSKR 130

- 45 A related DNA sequence was identified in *S. pyogenes* <SEQ ID 573> which encodes the amino acid sequence <SEQ ID 574>. Analysis of this protein sequence reveals the following:

Possible site: 56
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1865 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

- 55 An alignment of the GAS and GBS proteins is shown below:

Identities = 124/130 (95%), Positives = 129/130 (98%)
 Query: 1 MAQAQYAGTGRRKNVAVRVLVPGTGKITINKDVEEYIPHALLELVINQPFVAVTSTQGS 60

-249-

MAQAQYAGTGRKKHVARVRLVPGTSGKIT+NKKDVVEEYI PHADLRL+INQPFVATST+GS
 Sbjct: 1 MAQAQYAGTGRKKHVARVRLVPGTSGKITVNKKDVVEEYI PHADLRLINQPFVATSTEGS 60

Query: 61 YDVFNNVGGGYAGQSGAIRHGISRALLEVDGDFRDSLKRAGLLTRDARMVERKKPGLKK 120
 YDVFNNVGGGY GQSGAIRHGII+RALL+VDPDFRDSLKRAGLLTRDARMVERKKPGLKK
 Sbjct: 61 YDVFNNVGGGYGQSGAIRHGIRALLQVDPDFRDSLKRAGLLTRDARMVERKKPGLKK 120

Query: 121 ARKASQFSKR 130
 ARKASQFSKR
 Sbjct: 121 ARKASQFSKR 130

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 172

A DNA sequence (GBSx0178) was identified in *S.galactiae* <SEQ ID 575> which encodes the amino acid sequence <SEQ ID 576>. This protein is predicted to be recombinase (b1345). Analysis of this protein sequence reveals the following:

Possible site: 43
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1939 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AA29618 GB:AF217235 integrase-like protein [Staphylococcus aureus]
 Identities = 127/386 (32%), Positives = 205/386 (52%), Gaps = 18/386 (4%)

Query: 3 IHKYPKSKARNGYLYFVKIYMKD---SQRADHIKRGFTRKEAKQEARLIYKASGKI 59
 I XY K Y++ Y+ D ++ +RGF+T +EAK EA+L + QTEVQS 56

Sbjct: 2 IKKYKKKGSTAYMFVA--YLSTDPITGKQKRTTRRGFKTERAKIAEAKL---QTEVQS 56

Query: 60 EEFIKPTHKTYNEIFEKQYQAYQDMVPTTASRTLDMLHLPLVMGDLPIKISPLDQC 119
 F+ T+ E++E W + YQ+ V +T R L +F IL D+PI KI+ CQ
 Sbjct: 57 NGFLANDITTFPEYELMLBQYQNTVREGTYQRVLTLPDTALLENPDQVPIKKTVPYQC 116

Query: 120 NFITDKAKTFKHKKQIKSYTGKVFDFALMKLLKHNPMABEIMPKRKKTRIE--NYWTV 176
 I K + +IK I+ YT VP +A+ K++ NP A P++K++ + Y++
 Sbjct: 117 KVINQNNKYSYDIKAIKIRITYSNVFKYAVSLKIIVDNFPAHTKAPKKEAQDASTKYYS 176

Query: 177 QELQEFLAIVLQREPYKHYALFRLLAYSGLRKGELYALQWADIDFQETLSVDKSLGR-L 235
 EL++FL V E+ +YA+FR LA++G R+GBL AL W DIDE +T+S++K+ R
 Sbjct: 177 DELKQFLTFV--EDDPLYALIFRTLAFTGPRKRGELMALTNWIDIFTKQTISIKTKCARGA 234

Query: 236 DQQAIEKGRNDPFSVRKIKLDSSETISILQEWKISQKRAQLAVAPLSIQDQLFTYCTR 295
 + + + K S R I +D +T S+L+ W++ + E + S + +FT
 Sbjct: 235 NYKLQIQEPKTKSHTRTISIDDKTASVLKSWRTHQRVESLKYG-HNTSDKHQHVFTTVD 293

Query: 296 SGSEIPLHJADYINNVLSRIIRKGLKLSIPHGFRTEATLMIGIKGVDPVNTAKRLGHSS 355
 + +PL+ ++ N L I K+ K+I HGFRHTH +L+ E G+ RIGH
 Sbjct: 294 N---KPLYPEHCNKALDLICEKNSFKRIKRVHGFRRHTCGLLFPAGLSIQEVDRLRGDI 350

Query: 356 QWLTJYSHSTTTIGEORSVKQFADYL 381
 + T+D Y+H T D+ +EA Y+
 Sbjct: 351 KTTMDIYAHVTEKQRDQVADKAKYI 376

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 577> which encodes the amino acid sequence <SEQ ID 578>. Analysis of this protein sequence reveals the following:

-250-

Possible site: 39

>>> Seems to have no N-terminal signal sequence

----- Final Results -----

5 bacterial cytoplasm --- Certainty=0.3445(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

10 Identities = 109/386 (28%), Positives = 185/386 (47%), Gaps = 28/386 (7%)

Query: 3 IHKYPSKAKNGYL-YFVKIYNVDSQRADHIKRGF--RTRKEA--KDYEARLYLKASG 57
 I K K KNG + Y IY+ D +K RTRKE K A+ +L

15 Sbjct: 6 IMKITEHKIKGKTIIVYRASLYLGIDQMTGKRVKTSITGRTKEVNGKAKHAQDFDLNNGS 65

Query: 58 KLESEFKPTKHTYNEIFKQWYQAYQDMVEPTTASRTLDMPRIHLVPMGDLPIKISPLD 117
 ++ K KT+ E+ W + Y+ V+P T T+ HI+P +G++ + KI+ D

Sbjct: 66 TIKR--KVVIKTFKELSHLWLETYKLTVPQTYDATVTLARHIMPTLGNMKVKDITASD 123

20 Query: 118 CQNFITDKAKTFYNIKQIKSYTGKVPDFAIRMKLLKHNPMAEIIPIKPK--KRIENY 174
 Q I +K + N ++S KV + + L+ +N +II+P+++ K +++ +

Sbjct: 124 IQMLINRLSKTYVNYTAVRSVIRKVLQQGVILGLIDYNSARDIILPRKQPNNAKKVK-FI 182

25 Query: 175 TVQELQEFLAIVLQSEPPYKH-----ALFRLLAYSGLRKGEIYALKWADIDFQETLSV 228
 +L+ FL L+ +K Y L++LL +GLR GE AL+W DID + T++

Sbjct: 183 DPSDLKSPLE-HLETSCHKRYNLYFDVLYQLLSTGLRIGRACALEWGDIDLNTJTAI 241

Query: 229 DKSRLRLDGAIEKGTNDFSVRKIKLDSETISILQEWKISQKEKAQLAVAPLSIEQDF 288
 +K+ + K R I +D +T+ L+ + Q + Q+ + +

30 Sbjct: 242 NKTYNK--NLKFLSTAKTQSGRVISVDKKTLSLK---LVQMQRQLFNEVGARVSEV 295

Query: 289 LFTYCTRSGSIEPLHADIYINVLRSIIRKHGLKKISPHGFHHTATHMIEIGVDFVNTAK 348
 +F TR +R + L ++ G+++ + H #RTHA+L++ G+

35 Sbjct: 296 VFATPR-----KYPNASVRQSALDTCKCEAGIERPTFHAFRHTHASLLNNGISYKELQY 351

Query: 349 RLGRASSQMTLDITYSHSITTGEDRSV 374

Sbjct: 352 RLGRANISMVLDITYGHLSKGKEKAV 377

40 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 173

A DNA sequence (GBSx0179) was identified in *S. agalactiae* <SEQ ID 579> which encodes the amino acid sequence <SEQ ID 580>. Analysis of this protein sequence reveals the following:

45 Possible site: 61
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

50 bacterial cytoplasm --- Certainty=0.2477(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

55 >GP:AF63067 GB:AF158600 putative DNA binding protein
 [Streptococcus thermophilus bacteriophage Sfil1]
 Identities = 32/70 (45%), Positives = 46/70 (65%), Gaps = 3/70 (4%)

Query: 3 NRLKELRDNLTOADLAKVINTWOSQYKRYNGKTSLSIENSKILADFFGVSPFYLLGL 62

60 Sbjct: 2 NRL LR+ +T+ +LA+ I ++ K +G + +S +K IADFFGVGS+ YLLGL
 NRLYLRLRESKITYVELAEKIGVSKLTIVKLEHGTSKISREAKKLADFFGVGSYLLGL 61

-251-

Query: 63 D---NNSKIA 69

D N+S IA

Sbjct: 62 DTTTDSLIA 71

- 5 A related DNA sequence was identified in *S.pyogenes* <SEQ ID 581> which encodes the amino acid sequence <SEQ ID 582>. Analysis of this protein sequence reveals the following:

Possible site: 30

>>> Seems to have no N-terminal signal sequence

- 10 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.0680 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

- 15 An alignment of the GAS and GBS proteins is shown below:

Identities = 21/61 (34%), Positives = 34/61 (55%)

- Query: 1 MYNRLKELRKDNGLTQADLAKVINTNQSYGKYENGKTSLSIENSILADFFGVSIPIYL 60
 MY R++ LR+D TQ +A +++ + + Y K E G+ +L + + VSI YLL
 20 Sbjct: 1 MYPRIENLRREDNDFTKFVANLLSFSHANYAKIERGEVALMADVLVQFYKLYNVSIDYLL 60

Query: 61 G 61

G

Sbjct: 61 G 61

- 25 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 174

- A DNA sequence (GBSx0180) was identified in *S.agalactiae* <SEQ ID 583> which encodes the amino acid sequence <SEQ ID 584>. Analysis of this protein sequence reveals the following:

Possible site: 29

>>> Seems to have no N-terminal signal sequence

- 35 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.5278 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

- 40 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 175

- A DNA sequence (GBSx0181) was identified in *S.agalactiae* <SEQ ID 585> which encodes the amino acid sequence <SEQ ID 586>. Analysis of this protein sequence reveals the following:

Possible site: 60

>>> Seems to have no N-terminal signal sequence

- 50 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3762 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

-252-

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

5 Example 176

A DNA sequence (GBSx0182) was identified in *S.agalactiae* <SEQ ID 587> which encodes the amino acid sequence <SEQ ID 588>. Analysis of this protein sequence reveals the following:

```

Possible site: 59
>>> Seems to have no N-terminal signal sequence
10  INTEGRAL    Likelihood = -9.66    Transmembrane    40 - 56 ( 33 - 65)
    INTEGRAL    Likelihood = -5.79    Transmembrane    62 - 78 ( 59 - 81)

----- Final Results -----
15      bacterial membrane --- Certainty=0.4864(Affirmative) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

No corresponding DNA sequence was identified in *S.pyogenes*.

20 A related GBS gene <SEQ ID 8505> and protein <SEQ ID 8506> were also identified. Analysis of this protein sequence reveals the following:

```

Lipop: Possible site: -1    Crend: 7
McG: Discrim Score:    -16.96
GVH: Signal Score (-7.5): -2.95
Possible site: 57
25 >>> Seems to have no N-terminal signal sequence
ALON program    count: 2 value: -9.66 threshold: 0.0
    INTEGRAL    Likelihood = -9.66    Transmembrane    33 - 49 ( 26 - 58)
    INTEGRAL    Likelihood = -5.79    Transmembrane    55 - 71 ( 52 - 74)
30 PERIPHERAL    Likelihood = 10.87    14
    modified ALON score:    2.43

*** Reasoning Step: 3

35 ----- Final Results -----
      bacterial membrane --- Certainty=0.4864(Affirmative) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>
      bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

40 Example 177

A DNA sequence (GBSx0183) was identified in *S.agalactiae* <SEQ ID 589> which encodes the amino acid sequence <SEQ ID 590>. Analysis of this protein sequence reveals the following:

```

Possible site: 31
45 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.3276(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
50      bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

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No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 178

- 5 A DNA sequence (GBSx0184) was identified in *S.agalactiae* <SEQ ID 591> which encodes the amino acid sequence <SEQ ID 592>. Analysis of this protein sequence reveals the following:

Possible site: 44
>>> Seems to have no N-terminal signal sequence

10 ----- Final Results -----
bacterial cytoplasm --- Certainty=0.3482 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

- 15 A related GBS nucleic acid sequence <SEQ ID 9509> which encodes amino acid sequence <SEQ ID 9510> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:CAA30291 GB:X07371 RepM protein (AA 1 - 314) [Staphylococcus aureus]
20 Identities = 89/283 (31%), Positives = 145/283 (50%), Gaps = 26/283 (9%)
Query: 67 KVSLENTMTAYIKSKKYLAMQIETHLAITVCTAMTMFRATTGDGIHVLMMYDKQ 126
K+S D +T+ + + + I + + P+A + +++ YDK
25 Sbjct: 42 KLSFDAMTITVGNLWNSAKKLSDFMSLDPQIRLMDILQTKPKAKA---LQEKVYIEYKVK 98
Query: 127 KGQQRKARFPRLEFNPNKRLVDSEII---DTIIPFLDISISRADLAFDLFEVDCSEF- 182
K R R +EFNPNKL E++ II ++ED +R DLAFD FE D S++
30 Sbjct: 99 KADTWDRNRMRVEFNPNKL--THDEWMLKHNIIIDYMEDDGFRLDLAFD-FEDDLSDIY 155
Query: 183 -VLEKKGRPTATYKSPRSSTGTLETYKLGAPRSEKQVRLNKKKBQLQNGTDKDKDFAQSF 241
+ EK + T F +TG ETKY G+ S + +R+YNNKKE+ +N D D +++
35 Sbjct: 156 ALSEKALKRTV---FFGTGKAETKYFGSRDSNRFIRITYNKKKKERENA---DVDVSAE- 208
Query: 242 KMMWLEPQLRSRSIDEIFEVI-DTIIPK--FNKGLSIEITQIVYLIHIDRNINKKLH 298
H WR+E +L+ +D D I KP L+ L + +YL L+H+++ N +LH
40 Sbjct: 209 -HLNRVETELKRDMDVYNNCFNDLHILKPANWATLESLSKQAMVYL--LLHSESKWGLH 265
Query: 299 ENTRARVYKILETHQTSYDTYLGGLKDLKHERPRLNQLAYY 341
RN+R +YK+i++ + S D L+K L L+ Q+ ++
Sbjct: 266 RNSRRKYKQIIQ--EISSIDI/LDMKSTI/TDNEENLQKQINFW 306

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

45 Example 179

A DNA sequence (GBSx0185) was identified in *S.agalactiae* <SEQ ID 593> which encodes the amino acid sequence <SEQ ID 594>. Analysis of this protein sequence reveals the following:

Possible site: 32
>>> Seems to have no N-terminal signal sequence
10 INTEGRAL Likelihood = -15.55 Transmembrane 137 - 153 (133 - 157)

----- Final Results -----
bacterial membrane --- Certainty=0.7220 (Affirmative) < succ>

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Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 180

A DNA sequence (GBSx0186) was identified in *S. agalactiae* <SEQ ID 595> which encodes the amino acid sequence <SEQ ID 596>. Analysis of this protein sequence reveals the following:

```
Possible site: 18
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3406 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

The protein has homology with the following sequences in the GENPEPT database:

```
>GP:CAA33713 GB:X15669 pre protein (AA 1-494) [Streptococcus
agalactiae]
Identities = 171/402 (42%), Positives = 250/402 (61%), Gaps = 46/402 (11%)

Query: 1 MSYVARMAMKYKSGQLTAIYNHNERIFKHNSNKELDVSRSHLAYELINRDAQZTHKQIK 60
      1 MSY+VARM K K+G L + HNER+P+ HSNK+I+ +SHLAYELT+RD++ +Y KQIK
Sbjct: 1 MSYVARMAMKMKAGNLGGAFKHNERVFTSHNKIDINPSRSHLAYELTDRDRSVSYKQIK 60

Query: 61 EHINENRLSTRGVKRDAILCNEWILITSDKTFFDSLDKQTRFEPETAKDYPAEKYGDANI 120
      61 +++NEN++S R +RKDA+LC+EWILITSDK FF+ LDE+QTR FFETAK+YPAE YG++NI
Sbjct: 61 DYNNENKVSINRAIKRDVLCDEWILITSDKDFFEKLDEBQTRTFETAKNYFAENYGSNI 120

Query: 121 AYARVHLDSTFHHHGLGIVPMWQKGLSSKALFGNKEKLVAIQDELPKYLANEHGZNLQRGE 180
      121 AYA VHLDESTFHHH+G+VP +NKLSSKA-F ++B+L IQ++LE+Y++HGP L+RG+
Sbjct: 121 AYASVHLDSTFHHHGVVFFENGKLSKAMP-DREELKHIGQEDLPRYMSDHGVSLEGRG 179

Query: 181 IGSKKQILETAEFKKQRLNDNRKLADQHEELKALDDKISNV-NDTIA----- 229
      181 + S+ KH AEFK ++ +L +K+ +D++ + NDT A
Sbjct: 181 INBEAKHKTVAEFKRAMADME-LKEELLEKYHAPPPVDERTGELANDTEAPWHEKGFALM 238

Query: 230 -DKESRLNEL---EAKENAVGDLKQYLEKQSLAESIEDIKDIKLLQDRIQKEDLVQK 285
      230 ++S ++E E +W KQY+ E +L S ++D D E+L +
Sbjct: 239 FEVQSPFIRETTMQEKMDLR---KQYQERLKKLESSKKPLRD-----DLSHLELLDK 288

Query: 286 SFDGKLKDDEKTYNRLPQTASKHASSNAELKRDLVKAQSQNNHLSRELLNHRKTAEKQIK 343
      286 +K+D E AS+ AS +L KA+ N L NH K+ E I+
Sbjct: 289 KTKEYIKIDSE-----ASERAS-----KLSKAGYINTLE---NHSSKLEAKIE 329

Query: 346 LQGNRLKLDKVKMLDQVQKILNKLSLVNKEKAKFMPKQY 387
      346 ++ +K K + K LN+S + K F+ K QY
Sbjct: 330 CLESNQLQEKQATKLEAKALNESBLRELKPKKNFLGKRY 371
```

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 597> which encodes the amino acid sequence <SEQ ID 598>. Analysis of this protein sequence reveals the following:

```
LPXTG motif: 2025-2030

Possible site: 52

>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood = -10.08 Transmembrane 2034 -2050 (2030 -2053)
INTEGRAL Likelihood = -6.05 Transmembrane 21 - 37 ( 20 - 39)

----- Final Results -----
bacterial membrane --- Certainty=0.5034 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
```

The protein has homology with the following sequences in the databases:

>GP:AAD03320 GB:AF067776 extracellular matrix binding protein
[Abiotrophia defectiva]
5 Identities = 362/1396 (25%), Positives = 591/1396 (41%), Gaps = 87/1396 (6%)
Query: 636 KAEVLKKAHRAATQAKTEKDWLSPEQKKAQKEKAKARLDEGLKALKAAADSLTLKVTBE 695
+A+ + A +A AI+ + L+ E+K A+K K +A + L + A K T
10 Sbjct: 636 EAKNAVNNNAKAKGTAIDNNNNI.TAEKKAEEKAKVSAAKNATL.AGIDQA-----KTTAA 689
Query: 696 AFVDKEKNPDSIPNQHKAGTADQARKQALDSDLKVEQKELESIDNNLTITDEKAAAKKK 755
+ K I + + A A AI+ + ++ I LT +EK A +
Sbjct: 690 RNAVQNKGTTDINAVNPVPVAKPAANAALS---QAAVNKINEISQRPDLTREEQAQPMQD 746
15 Query: 756 VINDAYDVAKQTAMERANSYEDLTITIKDEFLS---NLPHKCGTPLKDQSDAIAEELKKQOE 812
V A D A A + + +T+ +D+ L+ NLP TP + +A+ + +
Sbjct: 747 VETARDAAMAVKASAAANNQAVTSARDQGLNAVNNLP----TPAA-KYPEALGHVRQAADA 801
20 Query: 813 TEKAIBGDKRTLPRDEKKEQIADSKERLKSQDKVQKDAKNADAIAKKAFEEBKVNIPQAHIP 872
+AI + L +E+ + + + KA +G I
Sbjct: 802 KRQAIRDNNNLTAESQADALRCQVDAACITAEAAINQHTINATLAKADSDGVKAI----- 855
Query: 873 GDLN---KDKELKLAELKQKADDTKEAIDVDKTLTDEDEKKEQKVTKAELEKAKTDVQNT 929
D+N + K L+Q A +AI+ + LT+ +EK + + L AKT V+
25 Sbjct: 856 NDINPQPRSKPAANQALEQVAAAKRQAINNNQLITDEEKAQAIQGVQALANAKTQVQAA 915
Query: 930 QTREELDKKVPPELKAIEQTHVKGNLBGKNAIETDLKKAHTETVAKINGDDTLDKATKE 989
+++ AI + + +G K +AI ++ A ++ G + L +
30 Sbjct: 916 NNNNGVNQAQTAGTATANNINPQGTQ---KAQAIKAI.EAAEQAKRLLEIQQRNDLITTEERN 972
Query: 990 AQVKEADKALAAAGKDAITKADDADKVSTAVTEHTPFIKAAHKTGLDKQAQVDANTALDKA 1049
+ + A KDA+ +A + V+ A +I+ + T +K DA A+D+A
Sbjct: 973 NALADLTAKAQAADKAVNQARNNITGVAGAKDNGVAQIQGINPTAVKFP---DARNAIDQA 1029
35 Query: 1050 AEKERGEINKDATLTITTEDKAKQLKEVSTALTKAKDNVKAATADAINDRNGVATIDAV 1109
A + E + LT E+KA +K+V+ A AK + A + +N+ A +G A I A +
Sbjct: 1030 ARDKAEFPQANTKLTDEECAAATKKVQCAARDAKAAIDRAGSNGDVNNAVNGKAAIQAI 1089
40 Query: 1110 HKAGQDLGARKSQVAKLSEAAKATDKISADPTLTSEKEBQSKAVDABLKAATEAVNA 1169
+ K A ++ AA A K I+A+ LT +EK K V+ E KA AV+A
Sbjct: 1090 KALDDSPQSAKD TAKAAIQNAADAQKAAITANNALTQEKEAAAIKQVEDEAAKAAQAVDA 1149
Query: 1170 ADTADKVDALGEGVTDIKNQHSKSGSIDARREAHGKRLRVAQETKKAIEKDPILTTEE 1229
+ + VD A +G+ I + ++ + +D+ A +K I D TLT EE
45 Sbjct: 1150 SRSKADVDRAKDGLQKISDV---PAVQPPKINAI.AAVDQAATDKGAVINDDITLTQEE 1205
Query: 1230 KAKQVKVDAAKERGMAKLNEAKDADALDKAYGSGVTDIKNQHSKSGDPVDA.RGLNENSKI 1289
K ++ VD + +N+A + +G I N ++ A + ++
50 Sbjct: 1206 KEAATKVDDEAAKARQAINDATSNADVAAKQAQGTQAINNPQT---PAKNAAKAAV 1261
Query: 1290 DEVAQATKDAITADPTLTITAEKETQRCQNDKEATKAKBELAKADADALFKAYGDGVTGI 1349
++ A A K AI D LT EK+ VD+E KA++ + A + +G +I
Sbjct: 1262 EQADAAKQKATKNDPNLTQREKDAALIKVQETNKARQAI.DAATTNADVTAKQNBGTQAI 1321
55 Query: 1350 KMQHKSQGLDVRKDEHKKALEAVAKRVTAHIBADPTLTPEVRQKQAEVQKELELATK 1409
++ K K + K A+ A+ + IR DP LT E + + KA+ V E A +
Sbjct: 1322 NAVFQTPKA---KTDARNAVTOAEDKKSATENDPNLTREEKDAAKAVDAEATKARNA 1377
60 Query: 1410 IAEAKDADEADKAYGDGVTAIKNAHVIGKGI.EARKDLAKKDLAEEAAKTALIIDKTLT 1469
I A D+ +G AI + + + +A+ D AK + +A + K I D LT
Sbjct: 1378 IDAATSNDDPTAKQNBGTQAI---NAVPTPKAKTD-AKNNVTQAADRIKDAIENDPNLT 1433
Query: 1470 DDQKBEQLLGVDTEYKANGIENIDAKDAAGVDKAYSIDGVRLAQYKGGQINIDRRNNAK 1529
+++ VD E K + IDAA A V ++G + I + + AK
65 Sbjct: 1434 REEKVAAKAVDAEAKKAKDAIDATSNADVTAKQNBGTQAI---NDVPTPTATKIDAK 1489
Query: 1530 EPLLKEADKVTKLINDPTLTEDQKVDQINKVBQAKLDAIKSVDDAQADAINDALGKI 1589

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		++AD I DP LT++K KV+ A ++D A++ + +G	
	Sbjct:	1490 NAVTQAADAKDAIEKDPNLTREBKDAAKAKVDAEAKKAKDAIDAATSNAVDTAQNNGT	1549
5	Query:	1590 EMINQYQHGDGVVRKATAKGOLSEAKVKALIAKDPTLTQADKDKQTPAVDAAKNLT	1649
		+ IN+ Q K AK ++A K I KDP LT+ +KD A VDA A	
	Sbjct:	1550 KAINDPVQ----TPTAKTDAKNAVQAADAKDAIEKDPNLTREBKDAAKAKVDAEAKKA	1605
10	Query:	1650 IAAVDKATTTGINSQELSGITAINKAYRPGBGVKARKEAAKADLSEAKVKALITNDP	1709
		A+D AT+ + + G AIN + K AK ++A K I ND	
	Sbjct:	1606 KDAIDAATSNAVDTAQKDGAKNAINAVPQ----TPTAKTDAKNAVQAADAKDAIENDA	1661
15	Query:	1710 TLTAKDK-AKQTEAVAKALKAIAAVDKATTAGINGQLSGKITAINKAYRPGBGVKAR	1768
		LT+ +K A + + A+A KA A+D AT+ + + +G AIN + K	
	Sbjct:	1662 NLTREBKDAAKAKVDAEATKAK-NAIDAATSNAVDTAQNNGTNAINDVPQ----TPTAK	1716
20	Query:	1769 EAAKADLREBAKVREALINDPFLTKADK-AKQTEAVAKALKAIAAVDKATTAGINGQL	1827
		AK +++ A + AI NDP LT+ +K A + + A+A KA A+D AT+ + +	
	Sbjct:	1717 TDAKNAVDAQATDKKSAIENDPALTREBKDAAKAKVDAEATKAK-NAIDAATSNAVDTAQ	1775
25	Query:	1828 LGKGTAINKAYRPGBGVEAHKAAKANLEKVKETKALISGDRYLSBTEKAVQKQAVEQ	1887
		G AIN + K AK +++ A + KA I D L+ EK K V+	
	Sbjct:	1776 KDAKNAINAVPQ----TPTAKTDAKNAVQAATDKKAAIENDPALTREBKDAAKAKVDA	1831
30	Query:	1888 ALAKALQVEBAKTEAVKLAENLGTVAIRSAVAGLAKDTQATLAALNEAKQKAIETAL	1947
		KA ++AA + V ++G KD A AK A A+	
	Sbjct:	1832 EAKKAKDAIDAATSNAVDTAQKDG-----KDAINAVPQTPTAKTDAKNAV	1878
35	Query:	1948 QAAETLAKITTTDAKLTEAQKAEQSENVSLAKLTAINTVRSQAISVKEAKDKGITAIR	2007
		QAR + + I D LT +K V KA + +A S A V + +G AI	
	Sbjct:	1879 QAATDKKSAIENDPALTREBKDAKVAKVDAAKAKDAIDAATSNAVDTAQKTEGTQAIN	1938
	Query:	2008 AAYVFNKAVAKSSAN	2023
		A VP AK+ + N	
	Sbjct:	1939 A--VPQTPTAKTDAK	1952
An alignment of the GAS and GBS proteins is shown below:			
	Identities = 77/396 (19%), Positives = 157/396 (39%), Gaps = 48/396 (12%)		
40	Query:	42 LNYELTNRDQAQNTYHKQIKHINENRLSTRGVRKDALILCNWITSDKTFFDSLDEKQTR	101
		L++B+ + ++QN K+I + + D E +I K +++ EK T	
	Sbjct:	338 LDPEILH-PRSQNVSKKISKQVEAKPFP-----DPASYKEKVIKALKPFVYEATSEKITN	389
45	Query:	102 EFF--ETAKDYFAEKYGDANIAYARVHLESTPHMLGIVPMNGKLSKALFG--NKEK	157
		+ E AKD +K + I+ G V + +A+ NK	
	Sbjct:	390 DAWLDENADLQKQKLEQYIIS-----GKVAISEGTFQKQIDAANYKYS	434
50	Query:	158 LWAIQDELKPYLNHSGFNLQRGISGKKKHLETAFFKQKRLDN--ADRKIADKHRL	214
		D LP + N + + + ++I K D N K E L	
	Sbjct:	435 SQTFDSLPSQYKQKQ--NKENBQKGRQLITQRLDLTKAIQSDNLTGKQTIQKEAL	492
55	Query:	215 KALDDKSNVNDITADKESRLKLEBAKWDVAVDGLQYE-----LEKQSLAESIE	264
		KA + I +VN T++ + + + + K + + K+Y EK+ A E	
	Sbjct:	493 KAFETGISVNVQTVSLQKQRLIVYKASEKSEKKEYPESTPAWHIPQKEKEVKAQKE	552
60	Query:	265 DIKDIELQLDRIQKEDLVKQSFQSGKLMDKSTYNNRLFQTASIKASNAFLKRLDVKAS	324
		++K + L+I ++ + + B + Q A K A + +I+ DL S	
	Sbjct:	553 ELKKLHDTTLEKINQDKMLTPDQQAQQLKQASVTFKQQAIRKSAQTLQLEHDLADYVS	612
65	Query:	325 QNHLSRELLNHRKTAEKNIKLSQENRKLKDKVMIDBOVK----IANKSLSVWKEKAKE	380
		+N + + + K+ K+ +++ KIK+ + + + + + KKKAK	
	Sbjct:	613 ENEGKNSIPDKYKSGNKDVLNKAIEVKLEAHEATKQATKEKPMWLSPEQKQKQKAKA	672
	Query:	381 FMPKQVYRETLISINTIANPIGLAKTAIRQKQNVDS	416
		+ + + + L + +L + + + + A +K D8	
	Sbjct:	673 RLDEGL--KALKAADSLSLIKVTEBAFVDEKKNFDS	706

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 181

A DNA sequence (GBSx0187) was identified in *S.agalactiae* <SEQ ID 599> which encodes the amino acid sequence <SEQ ID 600>. Analysis of this protein sequence reveals the following:

```
Possible site: 42
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.2544 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 182

A DNA sequence (GBSx0188) was identified in *S.agalactiae* <SEQ ID 601> which encodes the amino acid sequence <SEQ ID 602>. Analysis of this protein sequence reveals the following:

```
Possible site: 57
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.2045 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

The protein has no significant homology with any sequences in the GENPEPT database.

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 603> which encodes the amino acid sequence <SEQ ID 604>. Analysis of this protein sequence reveals the following:

```
Possible site: 57
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.2045 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

An alignment of the GAS and GBS proteins is shown below:

Identities = 102/111 (91%), Positives = 107/111 (95%)

```
Query: 1 MDYKKYQIIYAPDVLEKLRKRDYISQNYSTSGQHMEQIISDIEKLEVPFVGFDDE 60
+DYKKYQIIYAPDVLEKLRKRDYISQNYSTSGQ KMEQIISDIEKLEVPFVGFDDE
Sbjct: 1 LDYKKYQIIYAPDVLEKLRKRDYISQNYSTSGQKMEQIISDIEKLEVPFVGFDDE 60

Query: 61 KYGSKISKYHSTRGYTLKSDYIVLYHIEEENRVIDYLLPTSDYMLKLF 111
KYGSKI YHST+GYTLKSDYIVLYHIE EENR+VIDYLLPT+SDY+KLK
Sbjct: 61 KYGSKIYHSTKGYTLKSDYIVLYHIEEENRVIDYLLPTSDYMLKLF 111
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 183

A DNA sequence (GBSx0189) was identified in *S.galactiae* <SEQ ID 605> which encodes the amino acid sequence <SEQ ID 606>. Analysis of this protein sequence reveals the following:

```
Possible site: 13
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.1621(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
```

The protein has no significant homology with any sequences in the GENPEPT database.

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 607> which encodes the amino acid sequence <SEQ ID 608>. Analysis of this protein sequence reveals the following:

```
Possible site: 22
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.1596(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
```

An alignment of the GAS and GBS proteins is shown below:

```
Identities = 91/95 (95%), Positives = 93/95 (97%)

Query: 1 MYTAEDNRVATPQANKELVSEAMTVLNKKHNLTLSSALRLFLQNQVVVINEVDLLTTEELK 60
      M T +IDNRVATPQANKELVSEAMTVLNKKHNLTLSSALRLFLQNQVVVINEVDLLTTEELK
Sbjct: 1 MTTVKIDNRVATPQANKELVSEAMTVLNKKHNLTLSSALRLFLQNQVVVINEVDLLTTEELK 60

Query: 61 EKLFRKQPAEINIKNIEDVRQKFPYTSEVRSELGL 95
      EKLFRKQPAEINIKNIEDVRQKFPYTSEVR+ELGL
Sbjct: 61 EKLFRKQPAEINIKNIEDVRQKFPYTSEVRSELGL 95
```

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 184

A DNA sequence (GBSx0190) was identified in *S.galactiae* <SEQ ID 609> which encodes the amino acid sequence <SEQ ID 610>. Analysis of this protein sequence reveals the following:

```
Possible site: 56
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.4568(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
```

A related GBS nucleic acid sequence <SEQ ID 9513> which encodes amino acid sequence <SEQ ID 9514> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

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```

>GP:CAA6375 GB:X65276 ORF1 [Clostridium acetobutylicum]
Identities = 36/91 (39%), Positives = 51/91 (55%)

Query: 2 MSQIKLTPEELRISQAQKYTTGSQITDVLTVLTQEQAVIDENMDGTAFDSPEAQFNELSP 61
      M+QI +TPEEL+ AQ Y + I + + + I R W G A F ++ Q+N+L
Sbjct: 1 MAQISVTPEELKSAQAVYIQSKEIDQAIQKVNEMNSTIASEWKGQAPQAYLEQYNQLHQ 60

Query: 62 KITQFAQLLEDINQQLKRVADVVSQTDSDIA 92
      + QF LLE +NQQL K AD V + D+ A
Sbjct: 61 TVVQFENLLESVQQLNKYADTVAERDAQDA 91

```

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

15 Example 185

A DNA sequence (GBSx0191) was identified in *S.agalactiae* <SEQ ID 611> which encodes the amino acid sequence <SEQ ID 612>. Analysis of this protein sequence reveals the following:

```

Possible site: 21
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.4523 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

30 Example 186

A DNA sequence (GBSx0192) was identified in *S.agalactiae* <SEQ ID 613> which encodes the amino acid sequence <SEQ ID 614>. Analysis of this protein sequence reveals the following:

```

Possible site: 44
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.5339 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 187

A DNA sequence (GBSx0193) was identified in *S.agalactiae* <SEQ ID 615> which encodes the amino acid sequence <SEQ ID 616>. This protein is predicted to be chromosome assembly protein. Analysis of this protein sequence reveals the following:

```

5      Possible site: 61
      >>> Seems to have no N-terminal signal sequence

      ----- Final Results -----
10      bacterial cytoplasm --- Certainty=0.4620(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

- 15 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 188

A DNA sequence (GBSx0194) was identified in *S.agalactiae* <SEQ ID 617> which encodes the amino acid sequence <SEQ ID 618>. Analysis of this protein sequence reveals the following:

```

20      Possible site: 46
      >>> Seems to have no N-terminal signal sequence

      ----- Final Results -----
25      bacterial cytoplasm --- Certainty=0.4511(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

- 30 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 189

A DNA sequence (GBSx0195) was identified in *S.agalactiae* <SEQ ID 619> which encodes the amino acid sequence <SEQ ID 620>. Analysis of this protein sequence reveals the following:

```

35      Possible site: 20
      >>> Seems to have no N-terminal signal sequence

      ----- Final Results -----
40      bacterial cytoplasm --- Certainty=0.5249(Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
      bacterial outside --- Certainty=0.0000(Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

- 45 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 190

A DNA sequence (GBSx0196) was identified in *S.agalactiae* <SEQ ID 621> which encodes the amino acid sequence <SEQ ID 622>. Analysis of this protein sequence reveals the following:

```

Possible site: 14
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3542 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 9515> which encodes amino acid sequence <SEQ ID 9516> was also identified.

The protein has no significant homology with any sequences in the GENPEPT database.

15 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 191

20 A DNA sequence (GBSx0197) was identified in *S.agalactiae* <SEQ ID 623> which encodes the amino acid sequence <SEQ ID 624>. Analysis of this protein sequence reveals the following:

```

Possible site: 15
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3098 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

30 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 192

35 A DNA sequence (GBSx0198) was identified in *S.agalactiae* <SEQ ID 625> which encodes the amino acid sequence <SEQ ID 626>. This protein is predicted to be rgg protein. Analysis of this protein sequence reveals the following:

```

Possible site: 59
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3177 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

45 The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AAA26968 GB:M89776 rgg [Streptococcus gordonii]
Identities = 74/277 (26%), Positives = 142/277 (50%)

```

Query: 7 IFREFRLNRQPSLKQVASNELSVSGLSRFERGESDLSLTKFLGALEAIDLSEIFMDRVN 66
 I + R ++ SLK+VA++SV+QLSR+ERG S L++ F L + +S++EF +
 5 Sbjct: 10 ILKIITRESKNLKEVAAGDISVAQLSRVERGISLTVDSFYSCLRNMSVSLAEQVYVH 69

Query: 67 KVQKSDQISLMSQMAQYHYQRDVAGLEKMSVBEGLKDKSSDIRCRINIVLFRGMICEC 126
 Y++D + L +++ + ++ LE ++ E ++ +IN ++ R + C
 Sbjct: 70 NYREADDVLSQKLSFAQRSMNTVKLESILAGSEMAQEFPEPKINYKINTIVIRATITVSC 129

Query: 127 DSSRMSEEDLCFLSDYLFQDKSWESIDYILIGNLYRYNTRHICQLVKEVINQKEYYRD 186
 + ++S D+ FL+DYLF + W + L N + +E+IN+ ++Y +
 10 Sbjct: 130 NPOYQVSKODIEFLDYLFSEBVGRYELNLFMSVNLLETLETLETASEMINRTQFYNN 189

Query: 187 IYINRNVEATLNNVETLIERALAEATFFLEKVEALLNNERNAYHRIILLYEKGFPLAY 246
 + NR + LLNVV IE L+ A FL ++ E + Y R++ + Y R + Y
 15 Sbjct: 190 LPENRRRIIKGLNVLNVSACIENNHQLQVAMFNTIYDNTKIPETDLYDRVLKIKYKHALYSY 249

Query: 247 AKGDSRGISQMGAIQPCQAIGSKHVENFQHFNRV 283
 G+ ++Q + F+ + S +E F R+
 20 Sbjct: 250 KVGNPFAHHDIEQCLSTFEYLDSPGVARKKEQPERI 286

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 627> which encodes the amino acid sequence <SEQ ID 628>. Analysis of this protein sequence reveals the following:

Possible site: 29
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3792(Affirmative) < succ>
 30 bacterial membranes --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 79/275 (28%), Positives = 146/275 (52%), Gaps = 11/275 (4%)

Query: 9 REFRLNRQPSLKQVASNELSVSGLSRFERGESDLSLTKFLGALBAIDLSEIFMDRVN 66
 R R +Q S+ +A LS SQ+SRFERGES+++ ++ L L+ ++++I BF+ +K
 35 Sbjct: 15 RRLKKGQVSIISFLADEYLSKSGISRFERGESSEITCSRLNLLDKANTITDFSFVSHSKT 74

Query: 69 QKSDQISLMSQMAQYHYQRDVAGLEKMSVBEGLKDKSSDIRCRINIVLFRGMICEC 126
 + +L+SQ + + +++V L K++ + KD R + +LF DS
 40 Sbjct: 75 H-THFFTLISQARKCYAEKNVVKLLKLL---KDYAHKDYE--RIMKAILF-----SIDS 123

Query: 129 SRSMSEEDLCFLSDYLFQDKSWESIDYILIGNLYRYNTRHICQLVKEVINQKEYYRD 186
 S S+e+L L+DYLF+ + W + IL+GN R+ N + L KE++ Y
 45 Sbjct: 124 SIAPSGEETRLTLDYLFKVBQWGYEIIILGNCSEPMNTNITFLTKEMVASFAYSEQNK 183

Query: 189 TNRNVEATLNNVETLIERALAEATFFLEKVEALLNNERNAYHRIILLYEKGFPLAY 246
 TN+ +V +N + I+ E + + + K+ LL +E N Y + + LY G+ +
 50 Sbjct: 184 TNKMLVQLSINCIIISIDHSCFENSRYLINKIDLLRDLNIFYETKTVFYVHYGYKLG 243

Query: 249 GDSRGISQMGAIQPCQAIGSKHVENFQHFNRV 283
 + G + M+QA+ F+ +G +++EH+ ++
 Sbjct: 244 EEMSGEEDMKQALQIFKYLGEDSIYYSYKEHYRQI 278

55 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 193

A DNA sequence (GBSx0199) was identified in *S. galactiae* <SEQ ID 629> which encodes the amino acid sequence <SEQ ID 630>. This protein is predicted to be permease. Analysis of this protein sequence
 60 reveals the following:

Possible site: 15
 >>> Seems to have no N-terminal signal sequence

5	INTEGRAL	Likelihood = -8.07	Transmembrane	217 - 233 (215 - 238)
	INTEGRAL	Likelihood = -7.96	Transmembrane	163 - 179 (158 - 185)
	INTEGRAL	Likelihood = -7.75	Transmembrane	71 - 87 (69 - 91)
	INTEGRAL	Likelihood = -7.22	Transmembrane	369 - 385 (356 - 389)
	INTEGRAL	Likelihood = -5.15	Transmembrane	279 - 295 (275 - 299)
	INTEGRAL	Likelihood = -4.88	Transmembrane	252 - 268 (250 - 270)
10	INTEGRAL	Likelihood = -4.78	Transmembrane	140 - 156 (139 - 157)
	INTEGRAL	Likelihood = -3.56	Transmembrane	343 - 359 (340 - 367)
	INTEGRAL	Likelihood = -3.13	Transmembrane	40 - 56 (39 - 56)
	INTEGRAL	Likelihood = -2.28	Transmembrane	94 - 110 (92 - 112)

----- Final Results -----

15	bacterial membrane --- Certainty=0.4227(Affirmative) < succ>
	bacterial outside --- Certainty=0.0000(Not Clear) < succ>
	bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

20 >GP:AAD36408 GB:AE001788 permease, putative [Thermotoga maritima]
 Identities = 97/396 (24%), Positives = 194/396 (48%), Gaps = 15/396 (3%)

Query: 1 MNINGIKLLSSRAVSKLGEVFDYDGNSTWIASMGGLQKILGIYQIVELLVSVILNPFPG 60
 MN N + S VS + G Y + W+ S G + + G+ I L +I++PF G

25 Sbjct: 1 MNRNLLFASGSPFVSLIGRTIYQVLAWMLYSKTGSBYV-GLFMSSSFLFAIIVSFPAG 59

Query: 61 ALADRFQRRKILLITDAICAM--CFLLSPFGDDKVMVYGLIVANAILVANPSPFAY 117
 + DR RR +++ D + ++ FL+ + + + + L+ + ++V +F +FA

30 Sbjct: 60 TVVDRHSRRRMVMVMDILRGFLMFLMEXFSELIAL--LLIVTVLVSFDFSFNFAV 117

Query: 118 KSYIFIVDKADITTYANLETIVQIISVSSFVLGLFLINFGIRITIVDAITFLISFL 177
 S +F++V K +++ N+ + + + P LG L+ G+ +++++FLIS +

35 Sbjct: 118 DSIILFDLVRKENLVRANSLYRLNKLKSLGPAISGLLVVGLAGVILNLSFLISGI 177

Query: 178 FLYAIKRVQVSKQKQKVAINKALLADIADGFTYIKKEKEIMFFLIILALLNIFLAMPNYL 237
 F IKVE L K K +N+ DI Yi+ + I+ ++ A+N F + L

40 Sbjct: 178 FEMFTKVEEKHLKKVSKS--RNMWDIKSALLYTSRVFVLVTILVIAIMNFFFGSMHVL 235

Query: 238 LP-FTNSLLKTSAGAVTILSISGSIIGALIARKI--KSSINSMGLMVPSSLGVTVMG 294
 LP + L K+ Y T++S+ + G +I + I ++S+ ++ LV L V V

45 Sbjct: 236 LPEHVSGLKGSWYVYGTLMMSLSPGLIVTFLMATIRTRASVKTLGIANVGKGLAVVFA 295

Query: 295 FPLSLFELPIWIPYSGSFLPNSLI/MPNINHFFSQVQIRVDEAYMRGVMSTPIATIMFPI 354
 W+ ++ FL T+FN+ + +G+ + E G++ S I ++ +P+

50 Sbjct: 296 MTQNH---WLMFMYFLIGIPQTLFNINVTILQLAIPEMRGKIFSLISASVPSLLPV 351

Query: 355 GTLPMITPSFALNSVFIIGCAILGLGFSYSK 390
 F S ++ + I G+ S +

Sbjct: 352 SYQFPGLSSVATAHIFITTSNALIAGGVLSLQR 387

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 631> which encodes the amino acid sequence <SEQ ID 632>. Analysis of this protein sequence reveals the following:

Possible site: 45
 >>> Seems to have no N-terminal signal sequence

55	INTEGRAL	Likelihood = -8.17	Transmembrane	172 - 188 (161 - 194)
	INTEGRAL	Likelihood = -8.07	Transmembrane	220 - 236 (218 - 242)
	INTEGRAL	Likelihood = -7.22	Transmembrane	311 - 327 (303 - 329)
	INTEGRAL	Likelihood = -5.26	Transmembrane	98 - 114 (96 - 118)
	INTEGRAL	Likelihood = -4.99	Transmembrane	347 - 363 (342 - 370)
60	INTEGRAL	Likelihood = -4.62	Transmembrane	154 - 170 (151 - 171)
	INTEGRAL	Likelihood = -4.25	Transmembrane	284 - 300 (281 - 306)
	INTEGRAL	Likelihood = -3.66	Transmembrane	378 - 394 (378 - 396)
	INTEGRAL	Likelihood = -3.56	Transmembrane	74 - 90 (73 - 92)
65	INTEGRAL	Likelihood = -2.39	Transmembrane	50 - 66 (49 - 66)

----- Final Results -----

bacterial membrane --- Certainty=0.4270(Affirmative) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

5

The protein has homology with the following sequences in the databases:

>GP: AAD36408 GB: AB001788 permease, putative [Thermotoga maritima]
Identities = 85/345 (24%), Positives = 171/345 (48%), Gaps = 8/345 (2%)

10

Query: 40 SLSLVAVYQSLSSVIGVLPNLFPGGVADSPKRRKIIITINILCGTACLVSLPTKQMLV 99
S V ++ + ++ F G + D R + ++ + IL G + L + L
Sbjct: 36 SSEYVGLFMISPLFAIVSPFAGITVDRHSRRNMVMDIIRGLVPMYLFMYPSSELT 95

15

Query: 100 YAVIL-THVILAFMSAFSSPSYKAFTRKIVKDSISQINSLLETTSTVIKVIIVMVAIFL 158
A++L V+++ +F +P+ + ++V+K+++ + NSL + K+ P + L
Sbjct: 96 MALLLIVTLVLSVFDSPFNPAVDLSLLPDIVRKENLVRANSLYRLKKNLSKILGPAIGSLL 155

20

Query: 159 YKLGIGHGVLLDGLSPLTAALLISPLFVNDSEVVIKERVITREIFNDLKGPKYVSHK 218
K++G+ GV+L++ LSPIL+ + FI +K +K+ R ++ D+K Y+ S +
Sbjct: 156 LKVVGLAGVILINLSPLISGIPMPFIKV--EEKHLAKVSKRRMMQDIKSAIYIRSVR 213

25

Query: 219 SIFIITVLSALVNFPLAAYNLLLPYSINQMFGEISTGLYGTFLTAEAGGPGIAGLSGFVN 278
I + ++ A++NFF + ++LLP G+ S +YGT ++ + OG I L +
Sbjct: 214 FILVTILVIAINFTQGMHVLFPFVSKLKG-SBWVGTLMHSLPQGLIVTFMATIR 272

30

Query: 279 KELSSNRLLIPLSLSGMLMLAPFFYIMPHNAILALSPALFSLFSLIFNIQFPLVKQD 338
S L L L GL + + + M N ++ L +F ++FNI +L+Q
Sbjct: 273 TRASVITGLNLNGVGLAVFV----FAMTGNHWMFAMYLIGPITLPMINIVITLQLA 328

35

Query: 339 VNDNLFGRVPGIIFITITILFMPIGTGFFSVALNPNNSPNLFIIGS 383
+ + G++F+I ++ +P+ GFF + + ++FI S
Sbjct: 329 IPEEMRGKIFSLISAVSPSLLEPVSYGFFGFLSSVYATAHIFITTS 373

An alignment of the GAS and GBS proteins is shown below:

35

Identities = 136/379 (35%), Positives = 229/379 (59%), Gaps = 6/379 (1%)

40

Query: 8 LLSSRAVSKLQGVDPYDYNSTWIASNOGLQKILGIYQIVELLSVLNPNFGGALADRFP 67
L+ S+ + ++GDV +D+ N+T++A + ++ +YQ +E ++ ++ N FGG +AD P+
Sbjct: 11 LVYSKVIYTRIGDVMFDFANNITFLAGLNPASLSLVAVYQSLSSVIGVLPNLFPGGVADSPK 70

45

Query: 68 RRKILLITDAICAINCFLLSGEDDKVMVYGLIVANAILAVSNAPSSPAYKSYIPEVDK 127
R+KI++ T+ C C +LSP+ ++ +VY ++ N ILA +AFSSP+YK++ EIV K
Sbjct: 71 RKKIIITNTILCGTACLVSLPTKQMLVYAVILVILNVLAFMSAFSSPSYKAPTKEIVKK 130

50

Query: 128 ADIITYNANLETIVQIISVSSPVLGFLIPNFGIRITLIVDAITFLISPLFLYAIKVRV 187
I N+ LET +I V+ P++ ++ GI L++D ++FLI+ L + I
Sbjct: 131 DSISQINLSLETTSTVIKVIIVMVAIFLYKILGIGHGVLLDGLSPLIAALLISPLFVND 190

55

Query: 188 QLSKQEKVAIKNILADIADGPTYIKKEKIMFLLIAALLNTFLAMFNYPPLPPTNSLLK- 246
++ +BKV I+ I D+ GP Y+ K I +++AL+N FLA +N LLP++N +
Sbjct: 191 EVVIEKVTIREIFNDLKGPKYVSHKSIPIITVLSALVNFPLAAYNLLLPYSINQMFGE 250

Query: 247 -TSGAYATILSISAIGSIIGALLARKIKSSINSMKLVFSSLGVIIVMGPPS--LFLSLP 302
++G Y T L+ AIG IGA+++ + ++SM +L S G+++M P +P
Sbjct: 251 ISTGLYGTFLTAEAGGPGIAGLSGPNKELSSMRLLIPLSLSGMLMLAPFFYIMPHN 310

60

Query: 303 IWIPYSGSLFNLNLLTMFNHFPFSQVQIRVDEAYMGKVMSTIFTIAMIIMPGITLMTIF 362
I + S + LF+ L++FNI PFS VQ VD ++GRV IPTI I+FMPIGT P ++
Sbjct: 311 IIALSPA-LFSLFSLIFNIQFPLVQKVDNDNDFLGRVFGIIFPTITILFMPIGTGFFSVA 369

Query: 363 SPALSNVSFVIGCAAIL 381
++ + +IG I L
Sbjct: 370 INPNNSPNLFIIGSCITTL 388

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 194

A DNA sequence (GBSx0200) was identified in *S.galactiae* <SEQ ID 633> which encodes the amino acid sequence <SEQ ID 634>. This protein is predicted to be membrane permease OpuCD. Analysis of this protein sequence reveals the following:

```

Possible site: 46
>>> Seems to have an uncleavable N-term signal seq
10  INTEGRAL Likelihood = -5.68 Transmembrane 91 - 107 ( 88 - 110)
    INTEGRAL Likelihood = -4.30 Transmembrane 15 - 31 ( 9 - 37)
    INTEGRAL Likelihood = -3.72 Transmembrane 72 - 88 ( 72 - 88)
    INTEGRAL Likelihood = -3.19 Transmembrane 124 - 140 ( 123 - 142)

----- Final Results -----
15  bacterial membrane --- Certainty=0.3272 (Affirmative) < succ>
    bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
    bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 8509> which encodes amino acid sequence <SEQ ID 8510> was also identified. Analysis of this protein sequence reveals the following:

```

Lipop: Possible site: -1 Crend: 1
McG: Discrim Score: -10.69
QvH: Signal Score (-7.5): -3.79
    Possible site: 39
25  >>> Seems to have no N-terminal signal sequence
    ALOM program count: 5 value: -9.02 threshold: 0.0
    INTEGRAL Likelihood = -9.02 Transmembrane 35 - 51 ( 25 - 53)
    INTEGRAL Likelihood = -5.68 Transmembrane 151 - 167 ( 148 - 170)
    INTEGRAL Likelihood = -4.30 Transmembrane 75 - 91 ( 69 - 97)
30  INTEGRAL Likelihood = -3.72 Transmembrane 132 - 148 ( 132 - 148)
    INTEGRAL Likelihood = -3.19 Transmembrane 184 - 200 ( 183 - 202)
    PERIPHERAL Likelihood = 2.17 58
    modified ALOM score: 2.30

35  *** Reasoning Step: 3

----- Final Results -----
    bacterial membrane --- Certainty=0.4609 (Affirmative) < succ>
    bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
40  bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AAF91342 GB:AF249729 membrane permease OpuCD [Listeria monocytogenes]
Identities = 104/154 (67%), Positives = 133/154 (88%)

45  Query: 3 IANVIQTIPSLAMGISIMLGLGKIVVATVFLYSLLPIINITYTGIRNVDSLLIDAAK 62
    IAN+IQIPI+IAM+++ML+GLG TVV ++FLYSLLPI+ NIYTGIRNVD LL++ K
    Sbjct: 60 IANIITQIPALAMLAFLMLMIGLGTNTVVLFLYSLLPIILKNTYTGIRNVGGLLESOK 119

50  Query: 63 GGMGNIKQKLFMWVLEPLSISVIMAGLRNALVVAIGITAIGAFVGGGGLGDIIIRGINATN 122
    GGMGK Q L ++R+FL++SVIMAG+RNALV+AIG+ AIG FVG GGLGDII+RGINATN
    Sbjct: 120 AMGNIKVKVQLRLTLPALSVIMAGIRNALVIAIGVAAGITFVGAGGLGDIIIRGINATN 179

55  Query: 123 GGAITLAGSLPTALMAIPSLDLIAGIQRMLEPRK 156
    G AITLAG+PTA+MAI +D+LG ++R L P K
    Sbjct: 180 GTAITLAGAIPAVMAIADVLGHWERTLNPEVK 213

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 635> which encodes the amino acid sequence <SEQ ID 636>. Analysis of this protein sequence reveals the following:

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Possible site: 49
 >>> Seems to have no N-terminal signal sequence

5	INTEGRAL	Likelihood = -9.24	Transmembrane	39 - 55 (31 - 59)
	INTEGRAL	Likelihood = -7.17	Transmembrane	190 - 206 (188 - 211)
	INTEGRAL	Likelihood = -4.62	Transmembrane	93 - 109 (75 - 110)
	INTEGRAL	Likelihood = -3.66	Transmembrane	76 - 92 (75 - 92)
	INTEGRAL	Likelihood = -2.87	Transmembrane	221 - 237 (220 - 237)
	INTEGRAL	Likelihood = -2.44	Transmembrane	168 - 184 (165 - 184)

10 ----- Final Results -----
 bacterial membrane --- Certainty=0.4694(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

15 The protein has homology with the following sequences in the databases:
 >GP:AAD45530 GB:AP162656 choline transporter [Streptococcus pneumoniae]
 Identities = 344/508 (67%), Positives = 425/508 (82%), Gaps = 2/508 (0%)

20	Query: 13	MPSLFVTFQNRPNFNMIAAGHGLQISLLSMIALIGVPLAALLSRKRNDSIMLQVTVG 72
	M +L	TFQ+RF++NL AL +HLQ+SL+L++A+L+ +PLA L ++ +D +LQ+ G+
	Sbjct: 1	MTYLATFQDRPSDWLTALGQHLQSLTLTLAAILLATPLAVFLRYHEKLAADVWLQIAGI 60
	Query: 73	FQTIPSLALLGLFIFPLMIGITLFAVTALVVIYAFPILONTITGLANGIDPSLVEAGIARGM 132
	FQTI	PSLALLGLFIFPLMIGITLFA+TALVVIYAFPILONTITGL GIDF+L EAGIARGM
25	Sbjct: 61	FQTI
	PSLALLGLFIFPLMIGITLFAVTALVVIYAFPILONTITGLGIDFNIGIAGIARGM 120	
	Query: 133	TKWERLKTTFPIAMPVIMSGVSRSAVMIIGTATLASLIGAGGLGSFILLGIDRNANLI 192
	T+WERLK	FEIP+AMPVIMSG+RT+AV+IIGTATLA+LIGAGGLGSFILLGIDRNANLI
30	Sbjct: 121	TKWERLKTTFPIAMPVIMSGIRTAALVLIIGTATLAALIGAGGLGSFILLGIDRNANLI 180
	Query: 193	LIGAISSALLAIIPNSLLQYLEKASLRIMISFGITLALLASYTHMALSQFSKQKQTVV 252
	LIG+SSA+LAI	FN LL+ +EKA LR I F + L L SY+P L Q K K+ +
	Sbjct: 181	LIGAISSAVLAIAFNPFLKVMKAKLRITPFSGFALVALLGLSYSPALLVQ--KEKENLV 238
35	Query: 253	IAGKLGAEFDILINLYKELIEDQSDISVELKSNFGKTSFLYEALKSGDIDMYEFTGTIT 312
	IAGK+G	EP+IL N+YK LIB+ + + +K NFGKTSFLYEALK GSDID+YEFYGTIT
	Sbjct: 239	IAGKIGPEFELLANMYKLLIEBNTSMTATVKPNFGKTSFLYEALKSGDIDMYEFTGTIT 298
40	Query: 313	SSLLRDKPFLSNDPKOVYEDAKKGIAGDKLTLKPFAYQNTYAVAMFEKIAKEYOIRTI 372
	SLL+ P	+S++P+QVY+ A+ GIAKQD L LKP +YQNTYAVA+P+K++EY ++TI
	Sbjct: 299	ESLLQFSFKVSHPEQVYQVARDGIAQDHLVAILKPMSTQNTYAVAVPKIAQYEGYKTI 358
	Query: 373	SDLKANAUTLKGAGFTLEFKRADGYKMGSCYGLQLSVATMEPALRYQAIQSGDIQVTD 432
	SDLK	LKAGFTLEF DR DG K3+Q8 YGL L+VAT+EPALRYQAIQSGDIQVTD
45	Sbjct: 359	SDLKVBQGLKAGFTLEFNDREDGNGLQSMYGLNLVATTEPALRYQAIQSGDIQVTD 418
	Query: 433	YSTDAEITKYHLKVLKDDKQLFFPYQAPLMKTSLLTKHFLKGLIQAQKIKTEKMD 492
	YSTDAE+ +Y	L+VL+DDKQLFFPYQAPLMK +LL KHPEL+ +LN LAGKITE +M
50	Sbjct: 419	YSTDAELERYDLQVLEDDKQLFFPYQAPLMKTEALLKKHPELERYVLTLAGKITEBSMQ 478
	Query: 493	MYEYVSVKGADANKVARDYLLKTLGIQ 520
	+NY+V	V+G A +VA++L +GL+K
	Sbjct: 479	LNTQGVGVGSKAKQVAKFLQSGGLKK 506

55 An alignment of the GAS and GBS proteins is shown below:

Identities = 53/148 (35%), Positives = 93/148 (62%), Gaps = 1/148 (0%)

60	Query: 3	IANVIQTIPSLAMISIMLGLGLGIKTIVATVFLYSLIPITNTYTGIRNVDSLDLDAK 62
	+ V	QTIPSLA++ + + +G+G + V + +Y++ PI+ NT TG+ +D L++A
	Sbjct: 69	VTGVFQTIPSLAALLGLFIFPLMIGITLPAVTALVVIYAFPILONTITGLANGIDPSLVEAGI 128
	Query: 63	GMGNTKQRLFMVRLPLGISVIMAGLRNALVVAIGITAIGAFVGGGGLGDIIRGTATN 122
	GNIK +RL	E+P+++ VIM+G+R +V+ IG + + +G GGLG I+ G + N
65	Sbjct: 129	AFGNTKWERLKTTFPIAMPVIMSGVSRSAVMIIGTATLASLIGAGGLGSFILLGIDRN 188
	Query: 123	GGAILAGSLPTALMAIFSDLILOGIQ 150

-268-

+IL G++ +AL+AI + +L +++
 Sbjct: 189 AN-LILIGATISSALLAIFNLSLQYLK 215

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 195

A DNA sequence (GBSx0201) was identified in *S. agalactiae* <SEQ ID 637> which encodes the amino acid sequence <SEQ ID 638>. This protein is predicted to be choline transporter-related. Analysis of this protein sequence reveals the following:

10 Possible site: 44
 >>> May be a lipoprotein
 INTEGRAL Likelihood = -3.03 Transmembrane 306 - 322 (306 - 327)
 ----- Final Results -----
 15 bacterial membrane --- Certainty=0.2211 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 9517> which encodes amino acid sequence <SEQ ID 9518> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:CB15386 GB:Z99121 glycine betaine/carnitine/choline ABC
 transporter (osmoprotectant-binding protein) [Bacillus subtilis]
 Identities = 168/303 (55%), Positives = 224/303 (73%), Gaps = 1/303 (0%)
 25 Query: 2 LKSHPLQIFTLCLALLTISGQQLTVTKSGHTTIKVAAQSSTESSIMANITELIHHEL 61
 + K +L F L +L + GC L + TIK+ AQS TES I+AN+I +LI H+
 Sbjct: 1 MTKIKMLGAPALVFVML-LGGCSLPGLGASDDTIKIGAQSMTESEIVANMIAQLIEHDT 59
 30 Query: 62 GYNTLISNLGSSVTHQALLRGDADIAATRYTGTDTITGLGLKAVICPKKASKIVKTEF 121
 NT L+ NLGS+ V HQA+L GD DI+ATRY+GTD+T TLG +A KDPK+A IV+ EF
 Sbjct: 60 DLNTALVRNLGSSNVVQHQAMLGGDIDISATRYSGTDLTSLGKEARCPKALNIVQNEF 119
 35 Query: 122 QKRYNQTVPTYPYGFSDTYAFMVTKFARQNKITKIIDLKLLSTTMKAGVDSMMNRBEGD 181
 QKR++ W+ +YGF +TYAF VTK+FA + I +SDLKK ++ K GVD++W+ R+GDG
 Sbjct: 120 QKRFSYKMFDSYGFDMTYAFTVTKFAEKEHINTVSDLKKNASQYKLGVDNGLKRGD 179
 Query: 182 YTFDAKTYGFEFSHIYPMQIGLVYDAVSNKMGQSVLGYSTDGRISYDLEILRDDKFFFP 241
 Y F TYGFEP YPMQIGLVYDAV++ KM +VL YSTDORI +YDL+IL+DDK+FFP
 40 Sbjct: 180 YKGFVSTYGFEGTTPYPMQIGLVYDAVKNKMDAVLAYSTDORIKAYDLKILKDDKRRFFP 239
 Query: 242 PYEASMVNNSIIOKKPKLKKLLRLDGNKLNATKMNINVMVDDKLLPESVVAQFLEKN 301
 PY+ S V+ ++K+ P+L+ ++++L G+I+ +TMQ LMY VD KL EFSVAK+FLEK+
 45 Sbjct: 240 PYDCSPVIEPKVLEHPELEGVINKLIGQIDTMMQELNVEVDGLKRPESVVAKEFLEKH 299
 Query: 302 HYF 304
 HYF
 Sbjct: 300 HYF 302

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

A related GBS gene <SEQ ID 8511> and protein <SEQ ID 8512> were also identified. Analysis of this protein sequence reveals the following:

15 Lipop: Possible site: 22 Crend: 5
 McG: Discrim Score: 10.26
 GvH: Signal Score (-7.5): -4.19

-269-

```

Possible site: 44
>>> May be a lipoprotein
ALOM program count: 0 value: 8.65 threshold: 0.0
PERIPHERAL Likelihood = 8.65 66
modified ALOM score: -2.23

```

----- Final Results -----

```

bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

```

The protein has homology with the following sequences in the databases:

56.3/75.4% over 287aa

Bacillus subtilis

```

15      EGDJ109208|glycine betaine/carnitine/choline ABC Insert characterized
      SP102243|OPCC EAGSU    GLYCINE    BETAINE/CARNITINE/CHOLINE-BINDING    PROTEIN    PRECURSOR
      (OSMOPROTECTANT-BINDING
      PROTEIN). Insert characterized
      GP1263894|emb|CBR15386.1|Z99121 glycine betaine/carnitine/choline ABC transporter
20      (osmoprotectant-binding protein) Insert characterized
      PIR|E69670|R69670 glycine betaine/carnitine/choline ABC transporter (osmoprotec) opuCC -
      Insert characterized

```

ORF01181 (349 - 1212 of 1524)

25 EGBM109208|B3337615 302 of 303| glycine betaine/carnitine/choline ABC (Bacillus
subtilis) GP|032243|OPCC_EACSU GLYCINE BETAINE/CARNITINE/CHOLINE-BINDING PROTEIN PRECURSOR
(OSMOPROTECTANT-BINDING PROTEIN). GP|2635894|emb|CABL5386.1|299121 glycine
betaine/carnitine/choline ABC transporter (osmoprotectant-binding protein) (Bacillus
subtilis) PIR|B69670|B69670 glycine betaine/carnitine/choline ABC transporter (osmoprotec
30 opuCC - Bacillus subtilis
%Match = 33.5
%Identity = 56.2 %Similarity = 75.3
Matches = 162 Mismatches = 71 Conservative Sub.s = 55

35

VVFFLIVP+QCLIFIFSRYKSGSMKRINGVXQN+LXIITGNSNAQNKKRGGLDMKKSHFQLIPTLCALLTISQC
:::||
MTKIKWLGFALNPFVWLLGGCS
10 20

402 432 462 492 522 552 582 612

LTDTKKSHTTIKVAQSSTSESIWANNITELHHELGYNNTLISNGSSTVTHQALLREDADIAATRYTGDTITGLL

: |||: |||: |||: |||: |||: |||: |||: |||: |||: |||: |||: |||: |||: |||: |||: |||: |||:

YRLGASSTSTTGASSTSESIWANNITAGLIHEHTDNTALVQLGSSNYVQHALMGDDISDIATRYSGDTLSTLGR

45 40 50 60 70 80 90 100

642 672 702 732 762 792 822 852

KAVIQPKKAEASKIVKTEKQRKRNQTYPTPTGSDITAFVMTVEKPARQNKITKISDLKKISTTMAGDVGDSVAPRAGGGYTD

50 EAEKDKPKKALINIVONEPKRFSYKMFDSYGFQNTYATVTVTKFPAEKRIHINTVSLDKKASQYIKGLVDNAPLWKLRKGGYKGG

120 130 140 150 160 170 180

[illegible]

60
1122 1152 1182 1212 1242 1272 1302 1332
HRLOGKLNKLTNNYMDVDCILKEPSSVAQKFLDKHNYFGRGKMKQMNTFFQQFIYYFHNGSYLLSQFIHHPLLSVYG
||| :| :| || | | | | | | | | | | | | | | | | |
NKLIGQIDTETWQLNVYDVGKLEKPSVAKEFLDKHPD
280 290 300

65 SEQ ID 8512 (GBS23) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 14 (lane 8; MW 35kDa).

The GBS23-His fusion product was purified (Figure 194, lane 9) and used to immunise mice. The resulting antiserum was used for Western blot (Figure 251). These tests confirm that the protein is immunoreactive on GBS bacteria.

Example 196

- 5 A DNA sequence (GBSx0202) was identified in *S. galactiae* <SEQ ID 639> which encodes the amino acid sequence <SEQ ID 640>. This protein is predicted to be membrane permease OpuCB (opuBB). Analysis of this protein sequence reveals the following:

```

Possible site: 34
>>> Seems to have no N-terminal signal sequence
10  INTEGRAL    Likelihood = -9.66    Transmembrane    25 - 41 ( 18 - 45)
    INTEGRAL    Likelihood = -7.96    Transmembrane    182 - 198 ( 174 - 202)
    INTEGRAL    Likelihood = -4.83    Transmembrane    61 - 77 ( 57 - 95)
    INTEGRAL    Likelihood = -4.09    Transmembrane    78 - 94 ( 78 - 95)
15  INTEGRAL    Likelihood = -1.22    Transmembrane    134 - 150 ( 134 - 150)

----- Final Results -----
        bacterial membrane --- Certainty=0.4864 (Affirmative) < succ>
        bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
20  bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:AAF91340 GB:AF249729 membrane permease OpuCB [Listeria
monocytogenes]
25  Identities = 121/208 (58%), Positives = 160/208 (76%)

Query: 1  MYNFLSQYGMQLVKIWSQVYISFFAIALGIAIVELGVVLTFRFPKAKIIITAIASMLQT 60
+V F + G +LV+TW+ ++IS A+ LGIA+AVP G++LTR PKVA +I + S+LQT
Sbjct: 4  IVTFVQENGHNLLVQVQWHLFISLSAVILGIAVAVPTGILLTRSPKVNPFVIGVSVLQT 63

30  Query: 61  IPSLALLAIMIPLFGIGKIPAIIVALFIYSLPILRNITYIGMNNVNPTEKDCAGMGMKPI 120
+PSLA+LA +IP G+G +PAI+ALFIY+LLPILRNIT+IG+ V+ L+ +GMGM
Sbjct: 64  VPSLAAILAFIIPFLGVGTLPAILIALFIYALLPILRNITFIVRGVDKNLIESGRGMNIN 123

35  Query: 121  QSIFQVELFLATPIIMAGIRLSTYIVIAMATLASYGAGGLDLPISGLNLQSKLILGG 180
Q I V2+P + +IHAGIRLS +YVIAMATLASYGAGGLGD IF+GLNL++ LILGG
Sbjct: 124  QLVNVEIFNSIVIHAGIRLSAVYVIAMATLASYGAGGLGDIFNGNLNLRPDLLILGG 183

Query: 181  TIPVILSLIIDYLLGLLELATTPKTR 208
IPV IL+L++++ LG LE LTP+ R
40  Sbjct: 184  AIPVTILALVVEFALGKLEYITPKAIR 211

```

A related GBS gene <SEQ ID 8513> and protein <SEQ ID 8514> were also identified. Analysis of this protein sequence reveals the following:

```

Lipop: Possible site: -1  Crand: 0
McG: Discrim Score: -9.08
GVH: Signal Score (-7.5): -1.86
Possible site: 37
>>> Seems to have no N-terminal signal sequence
ALOW program count: 5 value: -8.60 threshold: 0.0
50  INTEGRAL    Likelihood = -8.60    Transmembrane    25 - 41 ( 18 - 45)
    INTEGRAL    Likelihood = -7.96    Transmembrane    182 - 198 ( 174 - 202)
    INTEGRAL    Likelihood = -4.83    Transmembrane    61 - 77 ( 57 - 95)
    INTEGRAL    Likelihood = -4.09    Transmembrane    78 - 94 ( 78 - 95)
    INTEGRAL    Likelihood = -1.22    Transmembrane    134 - 150 ( 134 - 150)
55  PERIPHERAL  Likelihood = 2.70      156
modified ALOW score: 2.22

```

*** Reasoning Step: 3

----- Final Results -----

bacterial membrane --- Certainty=0.4439(Affirmative) < succ>

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the databases:

ORF01825 (301 - 927 of 1233)

GP|9651976|gb|AAE91340.1|AF249729.2|AF249729(4 - 212 of 218) membrane perase OpuCB

{*Listeria monocytogenes*}

 $\%Match = 30.2$

%Identity = 57.9 %Similarity = 79.9

Matches = 121 Mismatches = 42 Conservative Sub.s = 46

117 147 177 207 237 267 297 327
STCF*YLKTY*FLCYGRRLT*KYC*AYFATWFKIRSSC*P*E*LKGHCYCIPS*YVIRYILGRY*NGGSIHVNFLSQYQ
: | : : |
MDAIVTFQENG
1.0

[illegible][illegible]

837 867 897 927 957 987 1017 1047
NLFQSKLILGGTPIVILLIIDIYLLGLLELATPRTTTREA*ICLNKRTFYRYLHFA*PS*RFLVVN*PILKSLVIPQL
[::] [::] [::] [::] [::] [::] [::] [::] [::] [::] [::] [::] [::] [::] [::] [::]
NLRYFDLIIGGAIPVTILALVVEFALGKLEYRITPKAIRREARBGGE

190 200 210

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 197

A DNA sequence (GBSx0203) was identified in *S. agalactiae* <SEQ ID 641> which encodes the amino acid sequence <SEQ ID 642>. Analysis of this protein sequence reveals the following:

Possible site: 46

```
>>> Seems to have no N-terminal signal sequence
```

----- Final Results -----

bacterial cytoplasm --- Certainty=0.3531(Affirmative) < succ

bacterial membrane --- Certainty=0.0000 (Not Clear) < succx

bacterial outside --- Certainty=0.0000 (Not Clear) < succ

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAF91339 GB:AF249729 ATPase OpuCA [Listeria monocytogenes]

Identities = 230/380 (60%), Positives = 298/380 (77%), Gaps = 4/380 (1%)

Query: 6 IIEYQNINKVY-GENVAVEDINLKIYPGDFVCFIGTSGSGKTTLMRMVNHMLKPTINGTLL 64
+++++++ K Y G AV D+ L I G+FVCFIG SG GKTT M+M+N +++PT G +

Query: 65 PKGKDISTNPIELRRRIGYVIQNLMPHMTIYENIVLPKLLKWSEERAKARELIK 124
KDI +P++LRR IGYVIQ IGLMPHMTI ENIVLPKLLKWSEE K+ +A+ELIK
Shift: 61 INDKDMAEDPVKLRISIGYVIOIGLMPHMTIRENIVLPKLLKWSEKKQERAKELIK 120

-272-

Query: 125 LVLPSEYLDRYPSELGGQQRIGVIRALAADQDIIMDEPFGALDPTIREGIDLVKS 184
 LV-LPSE+LDRYP ELGGQQRIGV+RALAA+Q+IIMDEPFGALDPTIR+ +Q+ K+
 Sbjct: 121 LVLDLPSEFLDRYPYELGGQQRIGVIRALAAEQMLILMDRPPGALDPTIRDSIQSEFKN 180

5 Query: 185 LQESMGKTTILVTHMDALKLATKIIVMDNGRQVQSGTNDLLHHPATSFVEQMGIER 244
 LQ+E+GKTII VTHMDRA+KLA +I+M +G+VQ TH+LL +EA SPVE IG+R
 Sbjct: 181 LQKELGKTTIIVTHMDALKLADRIIVMKDGEIVQPDTEILRNEANSFVEDFIGDR 240

10 Query: 245 LHAQADITVPVKQIMANPVVITAEKKTITAEITIMRQKVDLLVTEKGLI-OPIDLR 303
 L+ A+ D+T V QIM NPVETTA+K+L AIT+M+KRV+LLV D G + OPID+E
 Sbjct: 241 LIEAKPDVTQVQIMNTNFSITADKSLQAATVMKEKRVDTLLVVDGIVKSGPDVDEQ 300

15 Query: 304 LSSKYKDRVSDILGHTDPTVMEDDLRMTAEKILKGLKYAPVVDHNNKQIVTRAS 363
 + + + V DI+ FTV ED LLR+T +RLK G KY PVVD + L GIVTRAS
 Sbjct: 301 IDLNRRATATSMVDIIEKIVFIVYEDTILLRDTVQRILRGKYIIPVVDKRLVGIIVTRAS 360

Query: 364 LVDMILYDIINGDTE--TEDQ 381
 LVD++YD ING E TE+Q
 Sbjct: 361 LVDIVYDSINGTLEDAITNQ 380

20

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 643> which encodes the amino acid sequence <SEQ ID 644>. Analysis of this protein sequence reveals the following:

Possible site: 39
 >>> Seems to have no N-terminal signal sequence

25 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3619(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>

30

An alignment of the GAS and GBS proteins is shown below:

Identities = 102/237 (43%), Positives = 165/237 (69%), Gaps = 1/237 (0%)

35 Query: 6 LIIEYQNKVYGENVAVEDINLITYPDVFCITGSGSGKTLARMVAHMLKPTNITLLE 65
 +I + N+K +G+ +++ +I +F +G SSGSKTLL+M+N +P+G +L
 Sbjct: 1 MIFPNVSKTFQQTQVLQEQTFQINDREFFVLVGFSGSGSKTLLMKNCLIEPSSGDILL 60

Query: 66 KGDISTINPIELRRRIGVQINIGLPHMTIYENIVLPLKWSSEAKRAKARELKL 125
 + ++ E+R IGV+Q I L P+T+ ENI +P+ +WS E R K EL+
 40 Sbjct: 61 MNPQTELDLEMRLEIGVQLQIALFNLVAENTAIIPMKQWSAESTIKTEHLLDK 120

Query: 126 VELP-REYLDYPSLGGQQRIGVIRALAADQDIIMDEPFGALDPTIREGIDLVKS 184
 V LP ++YLDYPS+LGG+QORIG++RA+ + I-LMDEFP ALDPI+R+ +Q+L+ S
 45 Sbjct: 121 VGLPAKDYLDYPSLGGGQORIGVIRALISHPKILLMDEFPALDPISRKQLQELMLS 180

45 Query: 185 LQESMGKTTILVTHMDALKLATKIIVMDNGRQVQSGTNDLLHHPATSFVEQMG 241
 L+E TI+ VTHD+DEA+KL ++ ++ G+VQ P + HEA +FV +G
 Sbjct: 181 LHKEPDMTIVFVTHDIDEAIIKLDRAVAILNBSRIVQLDRPEMIKTHANAPVNLFG 237

50 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 198

A related DNA sequence (GBSx0212) was identified in *S.agalactiae* <SEQ ID 645> which encodes the amino acid sequence <SEQ ID 646>. Analysis of this protein sequence reveals the following:

55 Possible site: 24
 >>> Seems to have no N-terminal signal sequence

60 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.4736(Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000(Not Clear) < succ>

-273-

bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

- 5 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 199

A DNA sequence (GBSx0213) was identified in *S.agalactiae* <SEQ ID 647> which encodes the amino acid sequence <SEQ ID 648>. Analysis of this protein sequence reveals the following:

10 Possible site: 38
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -1.06 Transmembrane 18 - 34 (18 - 34)
 ----- Final Results -----
 15 bacterial membrane --- Certainty=0.1426 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

No corresponding DNA sequence was identified in *S.pyogenes*.

- 20 A related GBS gene <SEQ ID 8515> and protein <SEQ ID 8516> were also identified. Analysis of this protein sequence reveals the following:

Lipop: Possible site: 20 Crend: 5
 Sequence Pattern: CQMN
 SRCFLG: 0
 25 MoG: Length of UR: 19
 Peak Value of UR: 2.60
 Net Charge of CR: 3
 MoG: Discrim Score: 7.77
 GVH: Signal Score (-7.5): -4.89
 30 Possible site: 25
 >>> May be a lipoprotein
 Amino Acid Composition: calculated from 21
 ALOM program count: 0 value: 13.21 threshold: 0.0
 PERIPHERAL Likelihood = 13.21 115
 35 modified ALOM score: -3.14
 *** Reasoning Step: 3
 ----- Final Results -----
 40 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

45 ORF01527 (346 - 465 of 1095)
 EGAD|7398|7198(2 - 41 of 47) lysis protein for colicin e9 precursor {Escherichia coli}
 EGAD|41475|43808 lysis protein { } SP|P13344|LYS5_ECOLI LYSIS PROTEIN FOR COLICIN E5
 PRECURSOR. GP|40543|emb|CAA33861.1||X15857 lysis protein (AA 1-47) {Enterobacteriaceae}
 50 GP|144373|gb|AAA98053.1||M30445 colicin release protein {Plasmid ColE5-099}
 PIR|JQ0330|JQ0330 colicin E5 lysis protein precursor - Escherichia coli plasmid ColE5-099
 %Match = 3.7
 %Identity = 35.0 %Similarity = 52.5
 Matches = 14 Mismatches = 19 Conservative Sub.s = 7
 55 135 165 195 225 255 285 315 345
 YIYFFHCRIYIILININ*FN*GI*NIQMIFCLHVKTITIKIRENFVILKLL*CW*IIVNPIIYLYIKYIKIRKNMR

Example 202

A DNA sequence (GBSx0216) was identified in *S.agalactiae* <SEQ ID 653> which encodes the amino acid sequence <SEQ ID 654>. This protein is predicted to be lectin, alpha subunit precursor. Analysis of this protein sequence reveals the following:

```

5      Possible site: 47
      >>> Seems to have no N-terminal signal sequence

      ----- Final Results -----
10      bacterial cytoplasm --- Certainty=0.0653 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

- 15 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 203

A DNA sequence (GBSx0217) was identified in *S.agalactiae* <SEQ ID 655> which encodes the amino acid sequence <SEQ ID 656>. Analysis of this protein sequence reveals the following:

```

20      Possible site: 41
      >>> Seems to have no N-terminal signal sequence

      ----- Final Results -----
25      bacterial cytoplasm --- Certainty=0.6569 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

- 30 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 204

A DNA sequence (GBSx0218) was identified in *S.agalactiae* <SEQ ID 657> which encodes the amino acid sequence <SEQ ID 658>. Analysis of this protein sequence reveals the following:

```

35      Possible site: 27
      >>> Seems to have no N-terminal signal sequence

      ----- Final Results -----
40      bacterial cytoplasm --- Certainty=0.5736 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

- 45 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 205

A DNA sequence (GBSx0219) was identified in *S.agalactiae* <SEQ ID 659> which encodes the amino acid sequence <SEQ ID 660>. Analysis of this protein sequence reveals the following:

```

Possible site: 52
5  >>> Seems to have an uncleavable N-term signal seq
    INTEGRAL    Likelihood =-13.11    Transmembrane    146 - 162 ( 138 - 170)
    INTEGRAL    Likelihood =-12.90    Transmembrane    13 - 29 ( 9 - 32)
    INTEGRAL    Likelihood = -9.50    Transmembrane    108 - 124 ( 104 - 129)
    INTEGRAL    Likelihood = -7.75    Transmembrane    40 - 56 ( 33 - 61)
10  INTEGRAL    Likelihood = -6.64    Transmembrane    177 - 193 ( 170 - 195)
    INTEGRAL    Likelihood = -3.35    Transmembrane    77 - 93 ( 77 - 97)

----- Final Results -----
15  bacterial membrane --- Certainty=0.6243 (Affirmative) < succ>
    bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
    bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 8517> which encodes amino acid sequence <SEQ ID 8518> was also identified.

20 The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 206

25 A DNA sequence (GBSx0220) was identified in *S.agalactiae* <SEQ ID 661> which encodes the amino acid sequence <SEQ ID 662>. Analysis of this protein sequence reveals the following:

```

Possible site: 37
>>> Seems to have no N-terminal signal sequence

30 ----- Final Results -----
    bacterial cytoplasm --- Certainty=0.2374 (Affirmative) < succ>
    bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
    bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

35 The protein has homology with the following sequences in the GENPEPT database:
    >GP:AAB89623 GB:AE000990 repressor protein [Archaeoglobus
        fulgidus]
        Identities = 34/62 (54%), Positives = 46/62 (73%)

40 Query: 11 LKQVREDIGMTQQLAIRIGVRRRTIGHLENNKYNPSLEMAIKIVKIDMKIEDIFQLRK 70
    +K+ R MTQ+ELA R+GVRRRTT LE +YNPSL++A KI ++F+ KIEDIF +
    Sbjct: 5 IKEPRAKFNMTQQLAKRVGVRRTIVFLKGGKYNPSLKLAIKYARVNAKIEDITFIFDE 64

    Query: 71 ED 72
    E+
45 Sbjct: 65 EE 66

```

There is also homology to SEQ ID 412.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 207

A DNA sequence (GBSx0221) was identified in *S. agalactiae* <SEQ ID 663> which encodes the amino acid sequence <SEQ ID 664>. Analysis of this protein sequence reveals the following:

```

Possible site: 36
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.3794 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:CAB61817 GB:ALL33236 putative acetyl transferase [Streptomyces
coslicolor A3(2)]
Identities = 30/97 (30%), Positives = 52/97 (52%), Gaps = 1/97 (1%)

Query: 82  VGLNIVTLARADNMONGELGYVFHNOFWNGYAFESILALLNSTYERKLGPHHIEAQITPG 141
          VM ++ + Q GE+ Y+ H + W G E +LL+ +++ GH I A P
Sbjct: 72  VGMGDLHVRSHTRQ-Q-GBISYIVHFRVWGQGGIGTEIGRSLLSLGFDRWGLHRIATCDPR 130

Query: 142  NERSEKIVRLRLGLTYETTRIDKPSFENGKWDKTLIYSI 178
          N+ S +++ +LG+TYE + ++ W D L+SI
Sbjct: 131  NQASSRVLTKLGNTYGBRRHRTAMIRDGWRKSLVFSI 167

```

No corresponding DNA sequence was identified in *S. pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 208

A DNA sequence (GBSx0222) was identified in *S. agalactiae* <SEQ ID 665> which encodes the amino acid sequence <SEQ ID 666>. This protein is predicted to be p20 protein. Analysis of this protein sequence reveals the following:

```

Possible site: 44
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.1044 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:CAA30415 GB:X07542 P20 (AA 1-178) [Bacillus licheniformis]
Identities = 56/175 (32%), Positives = 94/175 (53%), Gaps = 6/175 (3%)

Query: 16  TVLTERLRQLQPVETLVNDLFPSDDSETVFMQRYKANTVEEAQVLLA---NVCMKSP 72
          T+ TERL L+ +EL + + ++ SD E YM V +A+ ++ ++ ++
Sbjct: 3  TLYTERLTLRKMELEDADVLQYNSDPVETRYMNTPTFTVSGARDMIQINDLSLEGQA 62

Query: 73  GIYAMIEKESQIMIGLIELEIRDEFS--AEFGYILNKNYNGKQYMEACSKLSIGFEHL 130
          +++I KE+ ++IG + D+ + AE GY L +N+ GKG+ +EA KL+ GF L
Sbjct: 63  NRFSIIVKETDEVIGTCGFNMIDQENGRKRGYDGLGRNHWKGFASEAVQKLDYGFSL 122

Query: 131  DLRIYARFDINNKKSGNVMRIGMKGBSLRLHLAKNPGKGEWKRATYYSILKEBY 185
          +L RI A+ + N S ++ + +KRG LR K KG +S+LK EY
Sbjct: 123  NLNRIRKAEVPENTPSIKLNLNLSFQKGLLGRDYRK-AKGRLLDVYMSILLKREY 176

```

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 209

A DNA sequence (GBSx0223) was identified in *S.agalactiae* <SEQ ID 669> which encodes the amino acid sequence <SEQ ID 670>. Analysis of this protein sequence reveals the following:

```
Possible site: 51
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
bacterial cytoplasm --- Certainty=0.5180(Affirmative) < succ>
bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
```

The protein has homology with the following sequences in the GENPEPT database:

```
>GP:CAAS7001 GB:Z46902 unknown [Saccharomyces cerevisiae]
Identities = 105/224 (46%), Positives = 148/224 (65%), Gaps = 3/224 (1%)

Query: 1  MGDVVENFTBGKIPKIDTNGKTVRIEKINPD-HFEDLFQVYGLSTEDSLTYSFSKFN 59
      +G VE +T P+ L G T R+E ++ +H +LF YE + TY+ F
Sbjct: 11  VGADVEGHTTAFPEKVVGLGNTCLREPLDRERHSELFSAYSEAG-QKLWYTLPAQFPT 69

Query: 60  SKIEFDVFFQTLKSEDPYLAIVDNNITGKVLGTFSLMRIDTNRVVENGWVYSSKLKQ 119
      + E+ F + L +++D AI++ T +GT L+RID N +E+G-VV+S +L++
Sbjct: 70  MLEEYLFKIKELNETKDVFFAIINKETERAVGTLCLIRIDEANGSLVGVTVVPSPLQK 129

Query: 120  TRIATEAQYLVMKYVFEELCYRREYKWCDSLNAPSNNSAKGLGTFPGTFRQAVVYKGN 179
      T IATEAQ+L-MKYVF+L VRYEYKWCDSLN PS +A RLGF +EGTFRQ VVYKGR
Sbjct: 130  TIATEAQFLMKYVFDLLQYRREYKWCDSLNAGPSRAAMRLOFKYBGTFRQVYVYKGR 189

Query: 180  RDTWYSIILDKEMPEKKTFRPEKWLDDSNFVNGYQIRSLSSIEQ 223
      RDT W+SI+DKEM + FE+WLD +NF NG Q R +++I +
Sbjct: 190  RDTQWFSIIDKELWIRIKTPEWLDKTNFE-N3KQKRGIAAIRE 232
```

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 210

A DNA sequence (GBSx0224) was identified in *S.agalactiae* <SEQ ID 671> which encodes the amino acid sequence <SEQ ID 672>. Analysis of this protein sequence reveals the following:

```
Possible site: 39
>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood =-12.15 Transmembrane 25 - 41 ( 20 - 49)

----- Final Results -----
bacterial membrane --- Certainty=0.5861(Affirmative) < succ>
bacterial outside --- Certainty=0.0000(Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>
```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

A related GBS gene <SEQ ID 8519> and protein <SEQ ID 8520> were also identified. Analysis of this protein sequence reveals the following:

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Lipop: Possible site: -1 Crend: 10
 McG: Discrim Score: -3.31
 GVH: Signal Score (-7.5): -4.44
 Possible site: 39
 >>> Seems to have no N-terminal signal sequence
 ALOW program count: 1 value: -12.15 threshold: 0.0
 INTEGRAL Likelihood = -12.15 Transmembrane 25 - 41 (20 - 49)
 PERIPHERAL Likelihood = -11.94 59
 modified ALOW score: 2.93

*** Reasoning Step: 3

----- Final Results -----

bacterial membrane --- Certainty=0.5861(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

SEQ ID 672 (GBS43) was expressed in *E.coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 5 (lane 4; MW 34kDa). It was also expressed in *E.coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 13 (lane 9; MW 58kDa) and in Figure 15 (lane 4; MW 59kDa).

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 211

A DNA sequence (GBSx0225) was identified in *S.agalactiae* <SEQ ID 673> which encodes the amino acid sequence <SEQ ID 674>. Analysis of this protein sequence reveals the following:

Possible site: 32
 >>> May be a lipoprotein

----- Final Results -----

bacterial membrane --- Certainty=0.0000(Not Clear) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 9519> which encodes amino acid sequence <SEQ ID 9520> was also identified.

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 212

A DNA sequence (GBSx0226) was identified in *S.agalactiae* <SEQ ID 675> which encodes the amino acid sequence <SEQ ID 676>. Analysis of this protein sequence reveals the following:

Possible site: 44
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -1.54 Transmembrane 165 - 181 (164 - 181)
 INTEGRAL Likelihood = -0.85 Transmembrane 67 - 83 (67 - 84)

----- Final Results -----

bacterial membrane --- Certainty=0.1617(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

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The protein has homology with the following sequences in the GENPEPT database:

```
>GP:CAA62211 GB:Z28353 similar to a B.subtilis gene (GB:
      BACHEMEHY_5) (Clostridium pasteurianum)
Identities = 40/185 (21%), Positives = 87/185 (46%), Gaps = 6/185 (3%)

Query: 18 MPKGGKQKVLISAIELFASQSFHGTSTAQLAKNAEVSQATIKYKYPETKDKLLVFLEILVQ 77
      M K K + SAI++F++ G++G + ++A NA V++ T+Y +F++K+++ +I+E V
Sbjct: 1 MNCTKDNIFYSALKVPSNNGYNGATWDEIASNAGVAKGTLTYHPFKSKBEIFKYIIEEGVN 60

Query: 78 TIGRFPTTELSTFTSKRELHFPVQDRPKPIERNNDLIKILMQELLINSETSTIPTKLIN 137
      + T E + + + I KN D K++ +L ++
Sbjct: 61 LMQNSIDEATDKREKTALEKLAVCRVQLNLTYNRDFFKVIASQLWQKELRQLELDIMR 120

Query: 138 STDPNITKIFNCLSEGNL---NMKBILRAVIGQFITFIQLY-ILNIKPENLEELKQI 193
      + +I + E S+ N + + A +G + + LY ++N ++N+ ++ +
Sbjct: 121 NYVHIEEFVKDAMEAGSIKKGNLFAVAYFLGTLCS--VSLYEVINAENDHINTIENL 178

Query: 194 EKQIL 198
      IL
Sbjct: 179 MNYIL 183
```

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 213

A DNA sequence (GBSx0227) was identified in *S.galactiae* <SEQ ID 677> which encodes the amino acid sequence <SEQ ID 678>. Analysis of this protein sequence reveals the following:

```
Possible site: 24
>>> Seems to have no N-terminal signal sequence

----- Final Results -----
      bacterial cytoplasm --- Certainty=0.2389 (Affirmative) < succ>
      bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
      bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
```

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 214

A DNA sequence (GBSx0228) was identified in *S.galactiae* <SEQ ID 679> which encodes the amino acid sequence <SEQ ID 680>. Analysis of this protein sequence reveals the following:

```
Possible site: 43
>>> Seems to have no N-terminal signal sequence

INTEGRAL Likelihood =-13.32 Transmembrane 341 - 357 ( 333 - 361)
INTEGRAL Likelihood =-10.93 Transmembrane 253 - 269 ( 238 - 277)
INTEGRAL Likelihood =-10.77 Transmembrane 172 - 188 ( 166 - 196)
INTEGRAL Likelihood =-8.01 Transmembrane 225 - 241 ( 215 - 251)
INTEGRAL Likelihood =-7.01 Transmembrane 21 - 37 ( 18 - 42)
INTEGRAL Likelihood =-2.66 Transmembrane 285 - 301 ( 283 - 301)

----- Final Results -----
```

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bacterial membrane --- Certainty=0.6328(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

5 The protein has homology with the following sequences in the GENPEPT database:

>GP:CAB42664 GB:AL049819 putative integral membrane protein
 [Streptomyces coelicolor A3(2)]
 Identities = 60/156 (38%), Positives = 101/156 (64%), Gaps = 1/156 (0%)

10 Query: 176 LMGMVFVFVFLISGMALLKERTSGTLDRLLATPVKRSDIVFGYMLSYGILAIQITIVIV 235
 L+G +FL++ +A L+ERTSGTL+RLLA P+ +D++ GY L++G LAI+Q+ +
 Sbjct: 77 LLGIFPLITMFLVTSIATLEERTSGTLERLLAMPLSGKDLIAGYALAFGALAIQVSALAT 136

Query: 236 LSTIWLDDIQVVGSGIFSIVIIINFILALVALSLGIIMSTLAKSEFQMMQFIPLIIMPQLFF 295
 +W L + V GS + ++V + AL+ +LG+ VS A SEFQ +QF+P +I PQL
 Sbjct: 137 GLAVWFLGLDVTGSPWLLLLVALLDALVGLAGLGFVSAPFASEFQAVFMPAVIFPQLLL 196

Query: 296 SGII-PLENMAWAQTVGKILFLSYSGDALTKIIMY 330
 G+ P +NM + V +LMS+Y+ D + ++ +
 20 Sbjct: 197 CGLFTRDRNMFALEAVSDVLFMSYAVDGMNEVLRH 232

There is also homology to a DNA sequence which was identified in *S.pyogenes* <SEQ ID 681> which encodes the amino acid sequence <SEQ ID 682>. Analysis of this protein sequence reveals the following:

Possible site: 39
 >>> Seems to have no N-terminal signal sequence

INTEGRAL	Likelihood = -11.41	Transmembrane	263 - 279 (246 - 284)
INTEGRAL	Likelihood = -7.70	Transmembrane	231 - 247 (224 - 258)
INTEGRAL	Likelihood = -4.99	Transmembrane	20 - 36 (18 - 39)
INTEGRAL	Likelihood = -3.72	Transmembrane	349 - 365 (345 - 368)
INTEGRAL	Likelihood = -3.45	Transmembrane	187 - 203 (182 - 204)

30 ----- Final Results -----

bacterial membrane --- Certainty=0.5564(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 35 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP:CAB12662 GB:Z99108 similar to ABC transporter (ATP-binding protein) [Bacillus subtilis]
 Identities = 92/369 (24%), Positives = 180/369 (47%), Gaps = 25/369 (6%)

40 Query: 12 IKRKKTSYVTFPILPILITLLALSLSFSNNQAKIGILDKNSQISKQFIAQLKKNKYD 71
 I +K +Y+ F P+L T + S+ N+++ ++ I+D++ ++ +S+ +I QLK +
 Sbjct: 15 IFKFPQNYLMPAAPALLITFVFGSMLSGNDCKVRLAIVDQDDTILSGHYIRQLKAHIDMY 74

45 Query: 72 IFTKIRKKBHIDHYLDKSLSEAVLITDKGFSKDKVLQKSKQKINIRSIANSEITBWQAQTN 131
 +F + + L+ K + ++ I + F ++ +GK +L R VK
 Sbjct: 75 VFENMSKSKASEKLQKKIAGIIVISRSPTQLKRGKHPGLIFRNGPELSEAPMVQYAE 134

50 Query: 132 YLENNYNIIGDVALGNEDTFNR-----ILQKNQQLNYDVKQVTLDRSRSKAVSST 182
 L NI A T +K+++ + V +TL+D+ S T
 Sbjct: 135 SALATLNIQVTAKTASQTAGENKAAKYKTVFAKHREDIVPAVTRQTLSDKKEGEASDT 194

55 Query: 183 TT---GFLIILMIGSTSVIYSGILADKSSQLYHRIMLSNLSRFR---YMLSYVVCVFA 235
 + GF ++ ++ + IL + ++ RL+ ++SR Y+LS+ +G++
 Sbjct: 195 ASRAAGFSLFVMIITMGAGTILFARKNGVNSKLLTASVSRATIGAYVLSFFVIGWIG 254

Query: 236 FTQIVIMLSLLKVPNISFFVPTSLLLIIFPLSLLAIGFGLLIGATTONSQOSSQLANL 295
 F I ++LS +F I++ P ++++++ LF L +G GL+I A + +Q NL
 60 Sbjct: 255 FGI---LLSTHMLFGINWGNPAIVLVLS-LFLITVVGIGLMIANVNTEPQQLAPGNL 310

Query: 296 IVMPTSLAGCLWFLSITPSTMQAIGKLLPQNWVLSALA-IPQSGTISQAWVYLLAAG 354
 V+ T M+G WE+ I P +MQ+I + LPQ W+S + I +G ++ +L + G
 Sbjct: 311 FVIATCNVSGMYWIDIRKPAQSIABFLPQKAMSGLTEIANGARVTD---ILGICG 366

Query: 355 TALALISFS 363
 LA ++
 Sbjct: 367 ILLAFAAIT 375

An alignment of the GAS and GBS proteins is shown below:

Identities = 92/375 (24%), Positives = 164/375 (43%), Gaps = 66/375 (17%)

Query: 11 IKELF----RDKRTLAMFLAPILIMFLMNMVMSANSNTVKIGITINVTQVSNLDNIK 66
 IK LF R K + FL PIL L+ + S ++N + KIG ++ + +S
 Sbjct: 5 IKTLFVKIKRKKTSYVFFFLMFLITL-LALSLSPSNNQAKIGILDKDQNSQISK----- 59

Query: 67 HIQVRSKFFMSAKKALKSNKIDALISEDNSYTVFYAMTDSKTTLT-RQAFKTAIVNTH 125
 +F+ LK MK + ++ K + Y S + LT + F V
 Sbjct: 59 -----QFAQ----LQNKKYDIFTKIKREHIDHYLQDKSLRAVLITDKGFSDEKVLQG 107

Query: 126 NSKELISQVKILANKPKLAQSLQTRSYKIEKYNY-----GNKNT-----GF 168
 S++L I + N ++ + ++ ++ Y+ E YH GN++T +
 Sbjct: 108 KSKQL----NIRSIANSBITWVKAQVNYLLENYNIIGDVALGNEDTFFNRILQKNQQLNY 163

Query: 169 FAIMIPIL-----NGFMVFFVPLISGM--ALLKERTSGTLDRLLATPVRKSD 214
 K + + GF++ + S + +L +++S RL+ + + R
 Sbjct: 164 DVKCVLTDRSRKAVSSTTTFGLLILMLGSGTSVITSGLADKSSQLYHRLMLSNLSR-- 221

Query: 215 IVPGLMST---GILAIQTVIVLSTIMLLDIQVGSIFSIVIINFILVALSGILM 271
 F YMLST G +A IVI+LS + + I ++I+ F+L+H+ G+L+
 Sbjct: 222 --FRYMLSTVCVGFVAPTYIQVIMLSLKVFNISFFVPTSLILLIIFLFSLLAIGFGLLI 279

Query: 272 STLAKSEFQMMQFIPLIIMPQLFSGII-PLENMAAQAQTVKILPLSYSGDALTKIMY 330
 + ++ Q Q LI+HP +G + FL S+ Q +GK+LP ++ A I
 Sbjct: 280 GAITQNSQSSQLANLIVMPTSMLAQCLWFLSITSTFMAIGKLLPQNWGLSATA-IPQS 338

Query: 331 GQGLPNVSSNLLVLL 345
 G L L L L+
 Sbjct: 339 GSTLSQAWFYLLALM 353

A further related DNA sequence was identified in *S. pyogenes* <SEQ ID 9081> which encodes the amino acid sequence <SEQ ID 9082>. Analysis of this protein sequence reveals the following:

Possible site: 38
 >>> Seems to have an uncleavable N-term signal seq
 INTEGRAL Likelihood = -12.52 Transmembrane 21 - 37 (17 - 43)
 INTEGRAL Likelihood = -10.30 Transmembrane 351 - 367 (346 - 371)
 INTEGRAL Likelihood = -5.36 Transmembrane 262 - 278 (260 - 285)
 INTEGRAL Likelihood = -2.60 Transmembrane 288 - 304 (288 - 305)
 INTEGRAL Likelihood = -1.81 Transmembrane 229 - 245 (229 - 246)

----- Final Results -----
 bacterial membrane --- Certainty=0.6010 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS sequences follows:

Score = 62.5 bits (149), Expect = 9e-12
 Identities = 72/382 (18%), Positives = 166/382 (42%), Gaps = 32/382 (8%)

Query: 1 MYFLHLKKESLQIFRNRITALIAWVIFILMIVILSPAFKSSSEYFATTVPKLTIRYQLEG 60
 M + + +K +FR++ L NM + FIL++ ++ F ++ NT' ++ + ++
 Sbjct: 1 MRIIATTEKVIKELPRDKRTLAMFLAPILIMFLMNMVMSANSNTVKIGITINVTQVSN 60

Query: 61 EKTIDYQRFAPLFLKVLKQLKHLKTFKNSLEKDRQVRSBGALTAVLKVKKKQTKIVITRN 120
 L+ H++ + ++ + +A++ + N++ V N
 Sbjct: 61 N-----LDNIKHTQVRGKFNSSAKKALKSNKIDALIS-EDNSYTVFYAN 105

Query: 121 INQQLNADLINMLVKNYVDNAKTYDSIALY-----PQQLAHIRKRSVDYVKKVSIQTSK 174

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+ L K V+ + + I+ + P+ ++ RS Y+K + +
 Subject: 106 TDSSKTTLTQAFKTAVMNISKELISQVKILANKIPKLAQSQTRES-KYKE---KAY 161
 Query: 175 GATSDADYA----ISMPTMITTPYSMMNMLVLSDRQQRITNRHLTGVSFPLVFGKLI 230
 5 G + ++A I M M+ P+ + + +L+R +R+ T V S +VFG +
 Subject: 162 GNKNTGPFARMPILMGWVFFVFLISGMALLKERTSGYDRLLATVVRSDIVRGYML 221
 Query: 231 GAMLATTVQLSLYIPTRFVLKVMWGTNEMWLGITASLVVLSVAIGIGLGISIKKEAPL 290
 10 + +Q + + + T +L + + + +I + L +++++GI + K+E +
 Subject: 222 SYGLIATIQITIVLSTINWLDIQVWSIFSVIIVNPLALVALSLGLMSFLAKSEPM 281
 Query: 291 TVASNTIPIPAFLGGSYVELITLHSSIIINQLSNISPIQVNDLSLYLIPGGQXNP-IPV 349
 II F G +EL + +S + I P+ + D+L +I QQ P +
 15 Subject: 282 MQFIPLIIMPQLFPFG-IIFLENN-ASWACTVGKILPLSYSGDALTKIIMYQGLNVS 339
 Query: 350 TLIVNISIGTTFIILALIGMRK 371
 L+V + I I + G+++
 Subject: 340 NLNVLFLIILTIANIPGLK 361
 20 Based on this analysis, it was predicted that these proteins could be useful antigens for vaccines or diagnostics.

Example 215

A DNA sequence (GBSx0229) was identified in *S. galactiae* <SEQ ID 683> which encodes the amino acid sequence <SEQ ID 684>. This protein is predicted to be CG1718 gene product (b0794). Analysis of this
 25 protein sequence reveals the following:

Possible site: 61
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -1.17 Transmembrane 118 - 134 (117 - 134)
 30 ----- Final Results -----
 bacterial membrane --- Certainty=0.1468 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

35 A related GBS nucleic acid sequence <SEQ ID 8521> which encodes amino acid sequence <SEQ ID 8522> was also identified. Analysis of this protein sequence reveals the following:

Lipop: Possible site: -1 Crend: 8
 MG: Discrim Score: -10.96
 GVH: Signal Score (-7.5): -4.84
 40 Possible site: 15
 >>> Seems to have no N-terminal signal sequence
 ALOM program count: 1 value: -1.17 threshold: 0.0
 INTEGRAL Likelihood = -1.17 Transmembrane 142 - 158 (141 - 158)
 PERIPHERAL Likelihood = 4.98 197
 45 modified ALOM score: 0.73
 *** Reasoning Step: 3
 ----- Final Results -----
 50 bacterial membrane --- Certainty=0.1468 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

55 >GP:AAF50837 GB:AE003568 CG1718 gene product [Drosophila melanogaster]
 Identities = 80/204 (39%), Positives = 123/204 (60%), Gaps = 3/204 (1%)
 Query: 7 EIIGLIPSGAGKSTLIKTLNMEKADKGTALV--LDTQMPDRTLNQIGYMAQSDALYE 64
 E GLG +GAGK+T K N G R+ G A V L + +I IGY Q DAL +

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Sbjct: 1394 ECFGLLVNGAGKTKTTFFKMTGDERISSGAATVQGLSLESNNMSIYKMGICPFDALLD 1453
 Query: 65 SLTGLENLFFGKMKGIQKTELKQIITHISKVDLENQLDKFVSGYSGQMKRRISLAL 124
 LTG E L F ++GQ++ ++Q ++K +DK YSGG KR+Ls AIA+
 Sbjct: 1454 DLTGREVLRIFCMLRGQVSRIRQLSDELAKSPGMKHIDKQTHAYSGGNRRKLSIAIV 1513
 Query: 125 LQNPVTVLIDDEPTVGIDPSLRKKIWOELINIKDEGHSIFITTHVMDAB-LTSKVALLR 183
 +G+P+V+ LDEPT G+D+ RR++W + I+D G SI +T+H N+E L ++++
 Sbjct: 1514 IGSPSVTLIDDEPTGMDPAARQLNVMVCRILKDSGKSIVLTHSHMEBCRALCTRLAMVN 1573
 Query: 184 GNIIAFDTPLHLKKQFMVSTIEEV 207
 G + HLK +F+ I ++
 Sbjct: 1574 GEFKICIGSTGLNFKSKGLILKI 1597
 Identities = 73/216 (33%), Positives = 128/216 (59%), Gaps = 9/216 (4%)
 Query: 1 MEVFKGEIIGLSPSGAGKSTLIKIMLGMEKADKGTALV--LDTPMORNINLQIGYMAQ 58
 M +F+ EI L+G +GAGK+T I + GM GTA++ D + +G Q
 Sbjct: 536 MNMFEDITVLIGHNGAGKTTTISMLTGMFPPTSTAILNGSDIRTNIEGARMISLIGCP 595
 Query: 59 SDALYSSTGLNLLFPFKMKGIQKTELKQIITHISKVDLENQLDKFVSGYSGQMKRR 118
 + L+ + + ++ PF +MKG++ ++Q+ K++LE++ + S SGKKR+L
 Sbjct: 596 HNVLFDEMSVSHIRFPFRMGLRGKAVGEVAKYLFMLELDEKANVASSKLGSGMKRL 655
 Query: 119 SLAIALGNPTVLIDDEPTVGIDPSLRKKIWOELINIKDEGHSIFITTHVMDAB-LTSK 177
 S+ AL G+ V++ DEP+ G+DPS RR++W +L+ + G ++ +T+H MDEA+ L +
 Sbjct: 656 SVCCALCGDQKVVLCDEPSSGMDPSARQLN-DLQGEKVGRTLLTTHPMDEADVIGDR 714
 Query: 178 VALLRGNIIAFDTPLHLKKQFN----VSTIEEV 208
 +A++ G + T LKKQ+ VS ++ +F
 Sbjct: 715 IAIMCDGLKQQTSPFLKKQGSQYRLVSGVQNL 750

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 685> which encodes the amino acid sequence <SEQ ID 686>. Analysis of this protein sequence reveals the following:

Possible site: 59
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.43 Transmembrane 49 - 65 (49 - 65)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.1171 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the databases:

>GP: CAB12660 GB:299108 similar to ABC transporter (ATP-binding
 protein) [Bacillus subtilis]
 Identities = 151/316 (47%), Positives = 202/316 (63%), Gaps = 18/316 (5%)
 Query: 4 VOLTIVVKSYYKNGKA-VNDVSLSTERMIVGLLGFNGAGKSTLINLILGLIFSLGKIT 62
 +Q N+ K+Y CHK V +S S++ G +GLFGAGKST I+I GL+P SG IT
 Sbjct: 2 LQRENKRAY--GKKTIVKGISPLKKGSGPALLGFNGAGKSTTISMGIVPHDSGNIT 59
 Query: 63 VLQGS-QKTIRKISSIGIVVQDIAYVVDLTAYHNVHLPGLSYGLKGAQLKQVLSLEF 121
 V G K K +IG VQ+LA+YP LTA+EN+ +G +YGL + KK+ + LE+
 Sbjct: 60 VGGYVIGKETAKAKQKIGIVPQRIALYPTLTAHNLMPWKGMYGLHDEAKRAEVL 119
 Query: 122 VGLHSQAKQFPSPQSGQMKRRINLACALVHSPKLLI+DEPTVGIDPQSRHILSLRLN 181
 VGL +AK PSQMKRR+NT AL+H P+L+I DEPTVGIDPQSRHILE++ LN
 Sbjct: 120 VGLTERAKDKIETPSGQMKRRINIGALMHKPELLINDEPTVGIDPQSRHILETVQLN 179
 Query: 182 KEGATVIYTHYHMEVEALGVIFDMHGVQIEBQPFELERKRYVANLANQIVITLDSR 241
 + G TVIYT+HYHMEVE L+D I I+D G+I G K +L R + Q+ V+ +
 Sbjct: 180 ETGMTVIYTHYHMEVEFLDRIIGIIDQGEIATGVKTDLCSRLGZTIQTVSGINEA 239
 Query: 242 HL----ELADKPSRLTEDGKMLKIDNSD-----MTSVHQLTCANTPSIRHNL 291
 L LA D++ R L LKID S +TS++ + T +I ++

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Sbjct: 240 FLVAIRSLAHVNDVTVHRE---LELKIDISAAHHSKVVTSLIAERATAHINILLSQVQEP 295

Query: 292 NLEEIFLHLTGKKLRD 307
NLE +FL+LTG+ LRD

5 Sbjct: 296 NLERLFLNLTGRTLRD 311

An alignment of the GAS and GBS proteins is shown below:

Identities = 81/211 (38%), Positives = 125/211 (58%), Gaps = 2/211 (0%)

10 Query: 1 MEVFKEIGLIGLPGSGACKSTLIKIMLGMEKADKGTALVL-DTQMPPDRNINLQIGYMAQS 59
++ G I GL+GP+GACKSTLI +L+ G VL +Q R I +QIGY+ Q
Sbjct: 25 L61EAGNLYGLGPGNGACKSTLINLILGLIPLSSGKITVLGQSQKTIKISQIGYVPOD 84

15 Query: 60 DALYESLTGLENLFFGFMKGIOKTELKQIITHISRVVDLENQLDKFV9SGSGMKRRLS 119
A+Y LT EN+ FG + G++ +L+Q+ + V L +Q +F S +SGSGMKRRLL+
Sbjct: 85 IAVYPDLTAYENVELFGSLYGLKGAQLKKQVLKSLFVGLHSQAKQFP9SGSGMKRRIN 144

20 Query: 120 IAIALLGNPTVLILDEPTVGIDPSLRKKIWQBLINIKDEGHSIFITTHVMDRAE-LTSKV 178
+A AI+ +P ++I DEPTVGIDP R I + + + EG ++ TTH M+E E L +
Sbjct: 145 IACALVHSPKLIIFDEPTVGIDPQSRNHILESIRLNKKGATVIYITTHYMEVEALCDYI 204

Query: 179 ALLLRGNI IAFDTPLHLKKQFNVSTIEVFL 209
++ G +I L+K++ + ++ +
25 Sbjct: 205 FINDHGQVIREGPKFELEKRYVANLANQIIV 235

SEQ ID 8522 (GBS391) was expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 74 (lane 7; MW 30kDa). It was also expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 83 (lane 4; MW 55kDa).

GBS391-GST was purified as shown in Figure 217, lane 3.

- 30 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 216

A DNA sequence (GBSx0230) was identified in *S. agalactiae* <SEQ ID 687> which encodes the amino acid sequence <SEQ ID 688>. Analysis of this protein sequence reveals the following:

35 Possible site: 13
>>> Seems to have no N-terminal signal sequence

---- Final Results ----

40 bacterial cytoplasm --- Certainty=0.6732 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S. pyogenes*.

- 45 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 217

A repeated DNA sequence (GBSx0231) was identified in *S. agalactiae* <SEQ ID 689> which encodes the amino acid sequence <SEQ ID 690>. This protein is predicted to be ISL2 protein. Analysis of this protein sequence reveals the following:

50

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Possible site: 58
 >>> Seems to have an uncleavable N-term signal seq

----- Final Results -----

- 5 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

- 10 >GP:CAC18596 GB:AJ278419 IS1381 transposase [Streptococcus pneumoniae]
 Identities = 111/129 (86%), Positives = 117/129 (90%)
- Query: 1 MKAQAIVTSQGRIVSLDIANYCHDMKLFKMSRRNIGQAAILADSGYQIMKMYSAQQT 60
 MK QAIVTSQGRIVSLDI VNYCHDMKLFKMSRRNIGQA KILADSGYQG+MK+Y QAQT
 15 Sbjct: 1 MKTQAIVTSQGRIVSLDITVNYCHDMKLFKMSRRNIGQAAGKILADSGYGLMKIYPAQQT 60
- Query: 61 PRKSKLKPLTLEDKTYNHTLSKERIKVENIFAKVTFKIPSTTYRNRKRPFGLRNMLIA 120
 RKSSKLKPLT+EDK NH LSKER KVENIFAKVTFK+FSITYR+ RKRPFGLRNML A
 20 Sbjct: 61 SRKSKLKPLTVEDKACNHALSKERSKVENIFAKVTFKIPSTTYRNRKRPFGLRNMLSA 120
- Query: 121 GMINRELGF 129
 G+IN ELGF
 Sbjct: 121 GIINHELGF 129

- 25 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 218

A repeated DNA sequence (GBSx0232) was identified in *S.agalactiae* <SEQ ID 691> which encodes the amino acid sequence <SEQ ID 692>. This protein is predicted to be ISL2 protein. Analysis of this protein sequence reveals the following:

Possible site: 41
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----

- 35 bacterial cytoplasm --- Certainty=0.3996 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

- 40 The protein has homology with the following sequences in the GENPEPT database:

- >GP:CAC18595 GB:AJ278419 IS1381 transposase [Streptococcus pneumoniae]
 Identities = 110/125 (88%), Positives = 119/125 (95%)
- 45 Query: 1 MNYEASKQLTDVRPKRIVGQRTTFEEMLAVLKTAYQKHAKGGRTPKLSLEDLLMATLQ 60
 MNYEASKQLTD RPKRIVGQRTTFEEMLAVLKTAYQ KHAKGR PKLSLEDLLMATLQ
 Sbjct: 1 MNYEASKQLTDARPKRIVGQRTTFEEMLAVLKTAYQLKHAQGRKPKLSLEDLLMATLQ 60
- Query: 61 YVREYRTYEQIADFGIHESNLIRRSQWVEVTLQSGPTISKTHLSAEDTVIVDATEYVKI 120
 Y+REYRTY+IADFG+HESNL+RRSQWVE TL+QSG TIS+T L\$+EDTV++DATEYVKI
 50 Sbjct: 61 YVREYRTYEEIADFGVHESNLLRRSQWVEVTLQSGVTISRTPLSSEDTVMIDATEYVKI 120
- Query: 121 NRPKK 125
 NRPKK
 Sbjct: 121 NRPKK 125

- 55 No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 219

A DNA sequence (GBSx0233) was identified in *S.agalactiae* <SEQ ID 693> which encodes the amino acid sequence <SEQ ID 694>. Analysis of this protein sequence reveals the following:

```

Possible site: 57
>>> Seems to have no N-terminal signal sequence
INTEGRAL Likelihood = -10.40 Transmembrane 130 - 146 ( 123 - 156)
INTEGRAL Likelihood = -7.86 Transmembrane 169 - 185 ( 167 - 191)
INTEGRAL Likelihood = -6.90 Transmembrane 100 - 116 ( 95 - 118)
INTEGRAL Likelihood = -5.52 Transmembrane 199 - 215 ( 189 - 216)

----- Final Results -----
bacterial membrane --- Certainty=0.5161 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

The protein has homology with the following sequences in the GENPEPT database:

```

>GP:BAB04126 GB:AP001508 unknown conserved protein in others
[Bacillus halodurans]
Identities = 47/207 (22%), Positives = 95/207 (45%), Gaps = 14/207 (6%)

Query: 7 LQKENTLLBGRIDNSNNQYTDIMIVYLGA-SISPYHQELIRNDIVNMLEAQERQASLV 65
      L K+N      + N + Y D+++Y+R A S S E + +++ LLEAQ + S
Sbjct: 6 LKKNNEKRKLLTEENLKVIEDLLLYIRLAHSSQETELLTELLDHLLEAQAKGSAK 65

Query: 66 SVFGEDRHDHFQINQVISTPKISKKE-TLQRMDLAILLITQIMIFLGQYLITLALQQSV 124
      +VFG++ + +++I PK+ KE L +L++ T+ ++F G Y + V
Sbjct: 66 AVFGDNPKQYADEIIGEIPMVTKERRGLFAYGLSMFFATV--LVFSGIYRMLRYVYFQV 123

Query: 125 PDLIPITLLDLVLFALFISIIAVKIADTIYATYNFDK----SKEKKYFFRYIFLLISLI 180
      + + + A+ +I ++ IA ++ + + K F +I + +I
Sbjct: 124 GEAVSEVYVGT--ALITTIASIVIAMMFVVFVQYFRMCSFRINKVFEPFILMLGMI 181

Query: 181 AYILIGKYYHLP---FINILPMIYLI 203
      + Y P I IP+++Y +
Sbjct: 182 FALFFALLYFTPNVGRMIEIPVYLYFV 208

```

No corresponding DNA sequence was identified in *S.pyogenes*.

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 220

A DNA sequence (GBSx0234) was identified in *S.agalactiae* <SEQ ID 695> which encodes the amino acid sequence <SEQ ID 696>. This protein is predicted to be minor extracellular protease epr precursor (epr).

Analysis of this protein sequence reveals the following:

```

Possible site: 31
>>> Seems to have an uncleavable N-term signal seq
INTEGRAL Likelihood = -10.72 Transmembrane 10 - 26 ( 5 - 33)

----- Final Results -----
bacterial membrane --- Certainty=0.5288 (Affirmative) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

```

A related GBS nucleic acid sequence <SEQ ID 8523> which encodes amino acid sequence <SEQ ID 8524> was also identified. Analysis of this protein sequence reveals the following:

```

5      Ldop Possible site: -1      Crend: 8
      MoG: Discrim Score:      12.11
      GvH: Signal Score (-7.5): -4.02
      Possible site: 29
      >>> Seems to have an uncleavable N-term signal seq
      ALOM program count: 1 value: -10.72 threshold: 0.0
10     INTEGRAL Likelihood = -10.72 Transmembrane 8 - 24 ( 5 - 33)
      PERIPHERAL Likelihood = 13.74 219
      modified ALOM score: 2.64

      *** Reasoning Step: 3

15     ----- Final Results -----
           bacterial membrane --- Certainty=0.5288 (Affirmative) < succ>
           bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
           bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

20     !GB:Z99123 extracellular serine protease [Bacillus s...

      >GP:CAB15866 GB:Z99123 extracellular serine protease [Bacillus subtilis]
      Identities = 44/150 (29%), Positives = 80/150 (53%), Gaps = 14/150 (9%)

25     Query: 37 QMDTVESSVNHVSDSQLTEAQDMLDKFEKKPSKLLKQVELANLKNSSKKKEALQKRFK 96
      ++D V+ S N ++A+D + K EK +++ + A+NKL N + K+ LQKR
      Sbjct: 428 RLDKVKQSYRN-----VKDAIDKVAKAKEYKTQQTVDTAQINKLFWGTDKKNLQKRLD 481

      Query: 97 KAKDKYLKDEADKKATKDATDLVEILQAPSEENVLKAAAVNKLTVKESKEALQKRIIT 156
      + K +Y+ A+K A D V E++ + +V A++A+ KL K +LQKR++
30     Sbjct: 482 QVK-RYI-----ASKQAKDKVAKAEKSKKTDVDSQAQSAIGKLPAASEKSTLQKRLNK 533

      Query: 157 VKTQYGLIGNQTFSSSVAEITQQTANPAS 186
      VK+ Q+ S++ ++T+ A S
35     Sbjct: 534 VKSTNLTKTAQQSVSAEKKSTDANAQAQS 563
      Identities = 39/124 (31%), Positives = 64/124 (51%), Gaps = 2/124 (1%)

      Query: 35 TTQMDTVESSVNHVSDSQLTEAQDMLDKFEKKPSKLLKQVELANLKNSSKKKEALQKRF 94
      +++ +++ +N V + L AQ + EKK ++ + A+N+L K ALQKR
40     Sbjct: 521 SSEKSTLQKRLNKVKSTNLTKTAQQSVSAEKKSTDANAQAQSAVNLQAGKDKTALQKR 580

      Query: 95 FKAKDKYLKDEADKKATKDATDLVEILQAPSEENVLKAAAVNKLTVKESKEALQKRI 154
      K K K EA K T A V+ E+ ++++ A++AVN+L K LQKR+
45     Sbjct: 581 LDKVKKVAAAEAKKVETAKAK--VKQAEKDKTKSKTSQAQSAVNLQAKSEKTKLQKRL 638

      Query: 155 DTVK 158
      + VK
      Sbjct: 639 NAVK 642

```

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 697> which encodes the amino acid sequence <SEQ ID 698>. Analysis of this protein sequence reveals the following:

```

55     Possible site: 41
      >>> Seems to have no N-terminal signal sequence
      INTEGRAL Likelihood = -4.99 Transmembrane 24 - 40 ( 23 - 43)

      ----- Final Results -----
           bacterial membrane --- Certainty=0.2996 (Affirmative) < succ>
           bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
           bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>
60

```

The protein has homology with the following sequences in the databases:

```

      >GP:CAB15866 GB:Z99123 extracellular serine protease [Bacillus subtilis]

```

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Identities = 43/130 (33%), Positives = 71/130 (54%), Gaps = 8/130 (6%)

Query: 41 GSHPTQDKVA---KHSKSAALLKAVKAVNADRLATAAAIQEAQKAVDKLAESSKKK 97
 G P + + K + + + K + + LK A + + V + A + + T A + A Q AV + L K
 5 Sbjct: 516 GKLPASSSEKTSIQKRLNKVKSTNLKTAQQSVSAEKKSTDANAQAQSAVNLQAGKDKT 575

Query: 98 TLQRLN-----VAKAKQEQEDATQAVKAAEETLNQNLKDIAQKAVNDLSNKGKKAALQ 152
 I Q + + L + V A A + + + A V K A E + + K A Q AVN L + K L Q
 10 Sbjct: 576 ALQKRLDKKKVAAAEAKKVETAKAVKKAEBKDKPKSKTSQAQSAVNLQASNEKTKLQ 635

Query: 153 SRLDAILPAK 162
 RL + A + P K
 Sbjct: 636 KRLNAVKKPK 645
 15 Identities = 31/105 (29%), Positives = 53/105 (49%), Gaps = 1/105 (0%)

Query: 54 SKSAASLLKAVKAVNADRLATAAAIQEAQKAVDKLAESSKKKTLQRLNAVAKAQRFQ 113
 + + + S A + AV A + + I + A + + + L S K L + L + + + + +
 Sbjct: 380 AQATDSAYAAAGQAVGAGQTKAQIDINKARELISQLPHSDAKTALHKKRLDKVQSYRVK 439

Query: 114 DAATQAVKAAEETLNQNLKDIAQKAVNDLSNKGKKAALQSLDAI 158
 DA + KA E + Q D A Q A + N L N K L Q RLD +
 20 Sbjct: 440 DAKDKVAKA-EKYTKQTVDTAQTINKLPNGTDKSKLQKRLDOV 483

An alignment of the GAS and GBS proteins is shown below:

25 Identities = 61/233 (26%), Positives = 115/233 (49%), Gaps = 13/233 (5%)

Query: 2 SMKIDKELLALIASIILLIPASVTFFLFKDHGTTQMDTVSSVNVHSDSQLTEAQDMLD 61
 SM +KE L + S + + + + + F H TQ + S + + S L + A + +
 30 Sbjct: 12 SMTEKQKEALYVMSVLITITLIGGSLIFGSHPTQDKVAKHSKS--AASLLKAVKAVN 69

Query: 62 KFEKKPSEKLLKDELALNKLNSSSKEALQKRFKAKDKYLKDEADKKATKATDULVEI 121
 + + + + + A + RL + SSK + LQ + + AK K + + A AT V
 Sbjct: 70 DADRLATAAAIQEAQKAVDKLAESSKKKTLQBLNAVAKAQEQEDA-----ATQAVKA 122

Query: 122 LEQAPSENVKAEAAVNLKLVKSEKALQKRIDTVTKQGLIGNQTPSSVAETTEQOT 181
 E + + + A + AVN L + K K ALQ R + D + + I + + P S E I +
 35 Sbjct: 123 AEETLNQNLKDIAQKAVNDLSNKGKKAALQSLDAILPAKPII-DEFFPQS-GEITINSY 180

Query: 182 ANPASQDTSSYYNQVAPTYE-QPQANPTPTFGVNHVTP-TPGTVGVBATNG 232
 P D S + + + FT + + + + VTF + + P P T + P + + G
 40 Sbjct: 181 WTPFFGVDSDTYDINSQSPTLDPSSESSASDVTVPQSPHDPPIFPQTSSEPSDSG 233

SEQ ID 8524 (GBS278) was expressed in *E. coli* as a His-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 52 (lane 6; MW 40kDa).

45 The GBS278-His fusion product was purified (Figure 206, lane 10) and used to immunise mice. The resulting antiserum was used for FACS (Figure 305), which confirmed that the protein is immunoreactive on GBS bacteria.

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

50 Example 221

A DNA sequence (GBSx0235) was identified in *S. agalactiae* <SEQ ID 699> which encodes the amino acid sequence <SEQ ID 700>. Analysis of this protein sequence reveals the following:

Possible site: 53
 >>> Seems to have no N-terminal signal sequence

55 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1466 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>

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bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

No corresponding DNA sequence was identified in *S.pyogenes*.

- 5 Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

Example 222

A DNA sequence (GBSx0236) was identified in *S.agalactiae* <SEQ ID 701> which encodes the amino acid sequence <SEQ ID 702>. This protein is predicted to be N-acetylglucosamine-6-phosphate deacetylase (nagA). Analysis of this protein sequence reveals the following:

- Possible site: 15
>>> Seems to have no N-terminal signal sequence
- 15 ----- Final Results -----
bacterial cytoplasm --- Certainty=0.4607 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 9297> which encodes amino acid sequence <SEQ ID 9298> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

- >GP:AAQ21688 GB:AY007718 N-acetylglucosamine-6-phosphate deacetylase
[Lactococcus lactis subsp. cremoris]
Identities = 113/178 (63%), Positives = 135/178 (75%)
- 25 Query: 131 GIYFBGPYFTEYKGAQNPFIYMNPNLEEFAMQKAANGILITKIALAPERBEGVEEFVSAI 190
GI+FBGP+FTEE KGAQNP YMR+ + E MQ+AA G++ KI LAPEREG E+F+
Sbjct: 1 GIFFBGPFFTEKKGAQNPFIYMRDANMWELEDWQEAHGMILKJIGLAPEREGSEDIFRKA 60
- 30 Query: 191 TKQGVTVLGHSHNGTYKBAKKAAGASVWVHAYNGMRGLTHEEPGMVGA VVNLNTYAE 250
T+ GV +ALGHSN TYK+ A V+AGASVWVH +NQM G+TH+EPGMVGA+ N ENTYAE
Sbjct: 61 TESGVVIALGHSHNATYKQAVAGVQAGASVWVHTTNGMSQMTQEPGMVGA VVNLNTYAE 120
- 35 Query: 251 LICDGHVDPVACDIILMTQKGNHIFVALITDCAAGGAPDGDYMLGELPVVVSNTARL 308
LICDGHV P A +I++ EG +HV LITD M A G PDG YMLGE V V +G A L
Sbjct: 121 LICDGHVPEAAETIVVMKGADHVLLITDSMRAGLPDGPVYMLGEYEVVRDGAARL 178

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 703> which encodes the amino acid sequence <SEQ ID 704>. Analysis of this protein sequence reveals the following:

- Possible site: 40
>>> Seems to have no N-terminal signal sequence
- 45 ----- Final Results -----
bacterial cytoplasm --- Certainty=0.3114 (Affirmative) < succ>
bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

- Identities = 227/300 (75%), Positives = 262/300 (86%)
- 50 Query: 9 MTKYIKADRFYFADHVKNGYLEIKDNHFGKWHENISQGEILDYSGYQIAPGLVDTHIH 68
MT Y+KAD F+Y V+ GYL + D FG+W E + +I+DY+GYQIAPGLVDTHIH
Sbjct: 1 MTCYLKADCFYYFTEVRPAGVLSHDGVFGWTEIVPADAQIIDYTGTYQIAPGLVDTHIH 60

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Query: 69 GFAGADVMDCDSEGLRMSAGLLSTGVTSFLPTTLTSDTKRLBEASKSVAAVAGKQGA 128
 G+AGADVMD ++GI +MS GLL+TGVTSFLPTTLTS ++LE+ S ++A+VA + +GAK
 Sbjct: 61 GYAGADVMDNSAQSIHQMSBGLLATGVTSFLPTTLTSTFBLQKRVSGT LASVADQVKGA 120

5 Query: 129 IQSIYFEGPYFTREYKGAQNPIYMRNPLEBPAQMQKAAGLITKIALAPERGVVEFVS 188
 IQSIYFEGPYFTREYKGAQNP YM+ P LEEF WQKAAGSLI KIALAPER+GV+EFVS
 Sbjct: 121 IQSIYFEGPYFTREYKGAQNPSYMKTPLEEFDAWQKAAGSLI KIALAPERDGVKEFVS 180

10 Query: 189 AITKQGVTVLALGHSNTYKKAQKAVKAGASVWVHAYNGMRGLTHREPGMVGAVNLNPTY 248
 A+TKQGVTVLALGHSNTY+BAK+AV+AGASVWVHAYNGMRGLTHREPGMVGAVNLNPTY
 Sbjct: 181 AVTKQGVTVLALGHSNTYQBAKEAVQAGASVWVHAYNGMRGLTHREPGMVGAVNLNPTY 240

Query: 249 AELICDGHVDPVACDILMTQKQHNIVALITDCMAAGAPDGDYMLGELPVVNGTARL 308
 AELICDGHV P+ACDILM QKQH+HVA+ITDCM AGG+PDGUY+LGE VVV+NGTARL
 15 Sbjct: 241 AELICDGHVSPACDILMQKQHHDVAMITDCMRAGGSPDGYLLGEFSPVVGNTARL 300

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 223

- 20 A DNA sequence (GBSx0237) was identified in *S. agalactiae* <SEQ ID 705> which encodes the amino acid sequence <SEQ ID 706>. Analysis of this protein sequence reveals the following:

Possible site: 25
 >>> Seems to have no N-terminal signal sequence

- 25 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3709 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

- 30 A related GBS nucleic acid sequence <SEQ ID 9307> which encodes amino acid sequence <SEQ ID 9308> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:CA16112 GB:Z99124 yyaQ [Bacillus subtilis]
 Identities = 40/110 (36%), Positives = 62/110 (56%), Gaps = 12/110 (10%)

35 Query: 121 IAKTFEDSVDPFAKHQYASVYVSG--KWYALLPFLKMKLENVPAQLSED---HVEVL 175
 +++ S DYP+ K+P YAS R+ KMY L+ + +P+L D H++L
 Sbjct: 11 VKEKYGTSPDPWKEKYFNYSLRHTSNKKWYGLINV-----LPEKGLDGHGIDIL 63

40 Query: 176 NIKVNPQDMILLQKQGIYPSYHMSKCTWVSLDNTLSDIRIFKLVSQS 225
 N+K P+ + L E I P YHM K+ W+SIVL+ T + EI+ L+ S
 Sbjct: 64 NLKCPFEISDRLENGENILPGYHMDKEHWISIVLERTDPEGEYNLISQS 113

- A related DNA sequence was identified in *S. pyogenes* <SEQ ID 707> which encodes the amino acid sequence <SEQ ID 708>. Analysis of this protein sequence reveals the following:

Possible site: 22
 >>> Seems to have no N-terminal signal sequence

- 50 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.2541 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

- 55 Identities = 114/247 (46%), Positives = 169/247 (68%), Gaps = 1/247 (0%)
- Query: 7 MSTESDPFRKKRPIFSSLEEFQFIKSDQYIYQTFMNDPKAITTISLDGKLAGKVIDS 66

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MS+ +D+ F ++ I L +GF K D Y Y + FM+ +F+ A + I G I +VID
 Sbjct: 1 MSLATDYFSRQTPIVEKIMAYGPEKRGYFYNERFWBGEFEAQLRIDEAGNIWDRVIDC 60

Query: 67 ALEEEYLPRAANYNGSVGVEVSAYMAILGDISDCKDLFTQGSRLAEKIAKTPE 126
 LEE+YLP+ A + G+ G+VR+Y+ +L +S +C + F QANRLA+ I K +
 Sbjct: 61 DLEEDYLPQQAAWQFTYTGQVRAAYLELLERLSVACFEATPFSQMQRNLAKHITKES 120

Query: 127 DSVDPYPAKHFOYASVYVSGKWWALLPPLKMGKLENVPAQLSEDEVEVLATIKVPMQDEI 186
 D +DYPF KHP A+YRV GQWY++F L KL+ +P +L EV+ +EVMP+
 Sbjct: 121 DPMDYPFKHPDLATYRVGGKWWYMIPLSLADKLDQIPERLVGQTEVMVIVKPKAPQ 180

Query: 187 LLQKGGIYPSYIMSKKWIVLINTLSDIIFKLVSDBSKLVSHNKKSN-SEPEFWIIP 245
 LLQ+GGIY+YIMSKK W+SI+LD+ ++D ++ LV+ SR+LV+ N SN + P+H+IP
 Sbjct: 181 LLQKGGIYPAYIMSKKNWISIILEDKVTYDDKLWTLVTSRQLVNPXLSHNPQDYNVPI 240

Query: 246 ANPKPYD 252
 AN K+YD
 Sbjct: 241 ANLKYD 247

20 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 224

A DNA sequence (GBSx0238) was identified in *S. agalactiae* <SEQ ID 709> which encodes the amino acid sequence <SEQ ID 710>. This protein is predicted to be transposase for insertion sequence element is905.

25 Analysis of this protein sequence reveals the following:

Possible site: 61
 >>> Seems to have no N-terminal signal sequence

30 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1824 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

A related GBS nucleic acid sequence <SEQ ID 9601> which encodes amino acid sequence <SEQ ID 9602> was also identified.

A related GBS nucleic acid sequence <SEQ ID 9595> which encodes amino acid sequence <SEQ ID 9596> was also identified.

The protein has homology with the following sequences in the GENPEPT database:

>GP:AAA25167 GB:L20851 transposase [Lactococcus lactis]
 Identities = 325/391 (83%), Positives = 365/391 (93%)

Query: 12 MTQFTTELLNPLAQQKQIDIEFFRSLETAMNDLLQVELSAPLGYEPYDKAGYNTGNSRNG 71
 MTQFTTELLNPLAQQKQIDIEFFR+LETAMNDLLQ ELSPLOYEPYDK GYN+GNSRNG
 Sbjct: 1 MTQFTTELLNPLAQQKQIDIEFFRTSLETAMNDLLQVELSAPLGYEPYDKVYNGNSRNG 60

Query: 72 AYTRRFETKYGVNVLIIIPDRNGEFSPALIPSYGRRDHLSEMVIKLYRTGVTTREISDI 131
 +Y+R+PETKY V L IPDRNG FSPAL+P+YGRD+HLEEMVTKLY+TGVTTREISDI
 Sbjct: 61 SYSRQFETKYGTGVLISLIPDRNGEFSPALIPYGRDDHLEEMVTKLYGTGVTTREISDI 120

Query: 132 IERMVGHYSPATVSNISKATQBNVASFHERSLEANTVYLLDGTYLPLRRGTYSKCEIH 191
 IERMVGHYSPAT+SNISKATQBNVA+ FHERSLEANTV+VL+LDGTYLPLRRGTYSKCEIH
 Sbjct: 121 IERMVGHYSPATISNISKATQBNVATFHERSLEANTVSVFLDGTYLPLRRGTYSKCEIH 180

Query: 192 IALGVTISYGHKAILGYDIAFNENNASWSLLERFKGQGVQVSVSVSDGFGNLDOLIOQA 251
 IALG+T G KA+LGY+IAFNENNASWS LL++ + QG+QGVSVV+DGF GL+Q+I QA
 Sbjct: 181 IALGVTISYGHKAVLYEIAFNENNASWSLLDQLQNGIQGVSVVTDGFKGLDQIIIOQA 240

Query: 252 FPMKQQRCLWHLGRNIASKVRADRALILBQFKITTRAINVRAKQALQSFINEKPHY 311

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+P+AKQQRCL+HI RN+ASKVVRADRA+ILBQFKTYRA N+E A QAL++PI EWKP Y
 Sbjet: 241 YPLAKQQRCLIHISRLNASKVVRADRAVILBQFKTYRAENLBMVAQALNFIAMKPKY 300
 Query: 312 KKVIEITLBSIHNLLIFVEFPHQWISYIYNLIESLNKRIKQTKKKVVPFNEESLERYL 371
 +KV+E+LE+ +NLL PY+FP+QIW SIYSTNLIESLNKRIKQTKKKV+PFNEE+LERYL
 Sbjet: 301 RKVMESLEWTDNLLTFYQFPYQIWSIYSTNLIESLNKRIKQTKKKVLPFNEEALERYL 360
 Query: 372 VTLFSDYNFKQCORIHKGFGQCTDTLESLEFD 402
 VTLF DYNFKQ QRIHKGFGQC DTLESLEFD
 Sbjet: 361 VTLFEDYNFKQSQRIHKGFGQCADTLESLEFD 391

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 711> which encodes the amino acid sequence <SEQ ID 712>. Analysis of this protein sequence reveals the following:

Possible site: 15
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3054 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below:

Identities = 111/128 (86%), Positives = 122/128 (94%)
 Query: 12 MTQPTTELLNFLAQQQIDIEFFRSSLETAMNDLLQVLSAPFLGYEPYDKAGYNTGNSRNG 71
 MTQPTTELLNFLAQQQIDIEFFRSSLE AMNDLLQVLSAPFLGYEPY+K GYNTGNSRNG
 Sbjet: 1 MTQPTTELLNFLAQQQIDIEFFRSSLEIAMDNDLLQVLSAPFLGYEPYKEGYNTGNSRNG 60
 Query: 72 AYTRRFETKYGVNLLIIPDRNKEFSPALPSYGRDRNHLDEMVKLYRTGVTTREISDI 131
 Y+R+FETKYGVNLLIIPDRNKEFSP L+PSY RR+HLEE+VKLY+TGVTTRREISDI
 Sbjet: 61 TYSRQFETKYGLVNLIIIPDRNKEFSPVLLPSYARRDHLLEEIVKLYTGVTTRREISDI 120
 Query: 132 IERNYGDH 139
 I+EMYG H
 Sbjet: 121 IKERNYGDH 128

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 225

A DNA sequence (GBSx0239) was identified in *S.agalactiae* <SEQ ID 713> which encodes the amino acid sequence <SEQ ID 714>. Analysis of this protein sequence reveals the following:

Possible site: 32
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -12.42 Transmembrane 268 - 284 (260 - 286)
 INTEGRAL Likelihood = -6.32 Transmembrane 232 - 248 (231 - 254)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.5967 (Affirmative) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AD40365 GB:AF036485 hypothetical protein [Plasmid pNZ4000]
 Identities = 69/283 (24%), Positives = 133/283 (46%), Gaps = 9/283 (3%)
 Query: 11 INVDDLSLQRRFP-LPSELLAYARDENESS-FVRDIEGHIALVYQLLDTQGHVDDVRHVP 68
 IN ++ + E++ + + ++ Y D +ES+ +V D+ L L D +R++
 Sbjet: 19 INAEERATLLEDQSGIDEDITEYVTQNDSESTNYVYDINEDDQLFIPLAPYALDKDALRYIT 78

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Query: 69 RVIPPTLFLKEDGLFVLAMHNNILVKKALNKV---EKVDSPKHLLSLVTPASQKQYFDV 125
 + P + L + LP N I V AL +V S +L + + +
 Sbjct: 79 Q--PFGMLHKGVLPTF-NQSGIPEVNTALYSALNPEVKSVDFALETFLTVVVSFPII 135

5 Query: 126 LDTISEEDKILINDLRKRPKNENLARIANLQSGTVHLMGTQKQPHMLDLQNIQBDKEN 185
 I++R+L L L++ S+L L+LQ L + N L L
 Sbjct: 136 SRAITKKRNYLDKMLNRKTKNSDLVSLSYLQQTITFLSSAVQTNLSLDRLPKTHFGVGA 195

10 Query: 186 TRNKMQLQDAILEAROLSNMCSLNSQVQELS-SYNNVLNNLNLNDVITLTITISIGISI 244
 +++ +D IE Q+ M + +QV + + N++RNLD + LTI S+ +++
 Sbjct: 196 DQKXILFEDVQIRGEQVQRMFEITQVDRIDETIANLNNLADTPKFKLTITWSLTWAV 255

Query: 245 IAMVTSFGYMNVLKLPDSVDVAVVLLIITITITIMSLVNYI 287
 +++ FYGMNVKLP + W+L + I+ ++ + + I++ +
 15 Sbjct: 256 PTIISGFGYMNVLKLPAGNQYAWMLTLOISVVVIVAMLMKIV 298

SEQ ID 714 (GBS422) was expressed in *E. coli* as a GST-fusion product. SDS-PAGE analysis of total cell extract is shown in Figure 172 (lane 7; MW 60kDa).

GBS422-GST was purified as shown in Figure 219, lane 12.

- 20 Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 226

A DNA sequence (GBSx0240) was identified in *S. galactiae* <SEQ ID 717> which encodes the amino acid sequence <SEQ ID 718>. Analysis of this protein sequence reveals the following:

25 Possible site: 45
 >>> Seems to have no N-terminal signal sequence

---- Final Results ----
 30 bacterial cytoplasm --- Certainty=0.0783 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

35 >GP:CB61731 GB:AL133220 putative oxidoreductase. [Streptomyces
 coelicolor A3(2)]
 Identities = 100/306 (32%), Positives = 152/306 (49%), Gaps = 3/306 (0%)

Query: 3 KRYGVVSTAKVAPRFIEGVRLANGVEVAVSSRTLESACAPANKYHLPKYDKLEDMLA 62
 KVR+G++T +A RF + + EVVAV+RT SA+ PA ++ +P+AY B +
 40 Sbjct: 8 KVRWGILATGGMPARFTADLVLEDAVAVASAKTFAERFGPIFRAYGNETLAR 67

Query: 63 DESIDVIVYATINQDHYKVAKALLAGKHVLVEKPTLTLYDQANELFALAESCNLFMEA 122
 DE +DV+VYAT + H A L AG+VL EKEPTL +A EL ALA +FLMEA
 45 Sbjct: 68 DEVDVVYVATPHEAERTAGLCLGAEGRVLCSEKFTINARAAELVALARENGVFLMEA 127

Query: 123 QKSVFIRMTOVVIKLLASGEIGEVISISSTTAYPN-IDHVIWFRLELGGGVVHPWAPYA 181
 P+ +K+L+A G IGEV S+ + R+ GGG + + Y
 Sbjct: 128 NMWYCNPLVRRILKELVADGATGEVRSIQADFLAGPFPANIRLEDPAQGGGALLDLGVYP 187

50 Query: 182 LSYLQYLFDATITHASGTATFPKQSDSOSKLLQLSNGVLVDIFLTIRNLNPHMI IYG 241
 +S+ Q L T + A + DQ+ IL N L I + P+ I G
 Sbjct: 188 VSPQALLGEP-TDVAARAVLSBEGVDLQTCALLSYGNDALASHCSITGTGTSPASBITG 246

55 Query: 242 TSKRLIIPH-FWKTTHAKLVNMTSARTIQVNVSDFEKAYHVSQMLEGQRVSHNIT 300
 +EKR+ +P+ F+ H L R + + D + H++ + R +D +P
 Sbjct: 247 SGRIDVWNGPFFFDHFLVIRTGRODQEPADGAPRESLRHEAEFVRMLRAGETESP 306

Query: 301 QNTLSG 306
 + L G

Subjct: 307 LVPLDG 312

Based on this analysis, it was predicted that this protein and its epitopes, could be useful antigens for vaccines or diagnostics.

5 Example 227

A DNA sequence (GBSx0241) was identified in *S.agalactiae* <SEQ ID 721> which encodes the amino acid sequence <SEQ ID 722>. This protein is predicted to be valyl-tRNA synthetase (valS). Analysis of this protein sequence reveals the following:

Possible site: 36
 >>> Seems to have no N-terminal signal sequence
 INTEGRAL Likelihood = -0.00 Transmembrane 794 - 810 (794 - 810)
 ----- Final Results -----
 bacterial membrane --- Certainty=0.1001(Affirmative) < succ>
 bacterial outside --- Certainty=0.0000(Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000(Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:AA57558 GB:L08854 valyl-tRNA synthetase [Lactobacillus casei]
 Identities = 543/881 (61%), Positives = 679/881 (76%), Gaps = 12/881 (1%)
 Query: 5 LSPKYNFAVEBGRYQWLDQDVFKPSGDTAKPYSIVPPNVTGKLHLGHAMDTTLQD 64
 L+PKY+ VEEGRY WLD+DVFKPSGD+AKPYSIVPPNVTGKLHLGHAMDTTLQD
 25 Subjct: 27 LAKPYDHKAVEBGRYQWLDQDVFKPSGDKAKPYSIVIPPNVTGKLHLGHAMDTTLQD 86
 Query: 65 IIRQKRMQGFDTLWLPQMCHAGIATQAKVZERLREQISRYDLGREKFLDKVWENKDEY 124
 I+IRQR++GFTLWLPQMCHAGIATQAKVE+LR++GISRYDLGREK+KVWENKDE+
 Subjct: 87 IVIRQKRIEGFTLWLPQMCHAGIATQAKVEAKLRKIGISRYDLGREK+VQKVWENKDEY 146
 30 Query: 125 AATIKSQWKGWGLSDVYSRERFTLDGLSKAVRKVPVLDLYNKGMIYGRFIIINNDPAART 184
 A TI QW KGLS+DYSRERFTLD+GL++AVR+VFVLDLYN+G IYSGRI+I+NDP ART
 Subjct: 147 AKTHGQWAKWGLSLDYSRERFTLDKGLNQAQVRRVFVLDLYNGLIYGRGIYVNDPQART 206
 35 Query: 185 ALSDIEVIHKDVEGAFYHDMVYLEDGSRALVATTRPETHFGDVAVAVNPEDARYKDLIG 244
 ALSDIEVIHKD +GAFYH+ Y DGS +E+ATTRPETM GD AVAV+P D RYK+AG
 Subjct: 207 ALSDIEVIHKDDKGAFFHVKYFPADGSGYIELATTRPETMGETAVAVHPSDERYKDMVG 266
 Query: 245 QNVILPIINKPIIVADSHADPEFGTGVVKITPAHDPNDFAVQGRHNLFPQVFNNDGTM 304
 +ILP+ N+ IPI+ D + DPEFGTG VKITPAHDPNDF VG RH+L ++N MNDDGTM
 40 Subjct: 267 TELIILPLANKPIIIEIDAVDFEFGTGAVKITPAHDPNDFQVGNRHDLKINTINDGTM 326
 Query: 305 NELADFNKMDRFEARKAVVAKLBSLNLVKKIKTTHSVGHSRITGVVPEPLSTQNFVK 364
 NE A ++ GMDRFEARKA+VA L+ G L+K++ HVGHSRITGV VE RLSTQNFVK
 45 Subjct: 327 NENAGKYQGMDFEARKAVVADLLKAGLLKVPFIVHSVHSRITGVVEARLSTQNFVK 386
 Query: 365 MDQLAKNAL-ANQDTEKVEFYPPRPMUTPMEMNENHDMVVISRQLWGHQIPAWNYN-VN 422
 M LA+ AI A Q++ KV F P RF T++ WMEN+HDMVISRQLWGHQIPAWNYN
 Subjct: 387 MKPLAEALKAQGEPPKKVIVPVRFEHTYLQWNIENHDMVVISRQLWGHQIPAWNYNQT 446
 50 Query: 423 GEMVVGSDAPEG-DGNTQDEVDLWFWFSALWPFSTMGWPFTEADPKRYFPPTSLTVGY 481
 GE YVG +AP+ + W QD DVLWFWFSALWPFSTMGWPF+T+A D+KRY+PT TLTVGY
 Subjct: 447 GETYVGMKAPQDIENKQDQDVLWFWFSALWPFSTMGWPFNTADPYKRYPTTLTVGY 506
 55 Query: 482 DIIPFWWSRMIFQSLEFTRQPPSNVLIHGLIRDEBGRMSKSLGNIGIDPMNVIEKYGAD 541
 DII FWW+RMIFQ L FT ++FF LIHGL+RDE+GRMSKSLGNIGIDPMNVIEKYGAD
 Subjct: 507 DIIPFWWSRMIFQGLHPTHGRPFQYTLIHGLMRDEBGRMSKSLGNIGIDPMNVIEKYGAD 566
 Query: 542 ALRWFLNSGAPQDQVRSYKONDAWNFINKIWNISRYLWNNIGLITDQARENVEKVV 601
 ALRWFL G+ PQD RPSY+++++WNFINKIWNISRYLWNNIGLITDQARENVEKVV
 60 Subjct: 567 ALRWFLITGNKPGQDTRPSYKONDAWNFINKIWNISRYLWNNIGLITDQARENVEKVV 620

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Query: 602 MSQGVNVDRIWILHNLNETVQKVTENFDKPEFGVAGHILYNFIWEEFANWVVELTKVLY 661
 +++D+W+ LNET+ +V + +FEFG G LYNF W A+WYVE++KEVLY
 Sbjct: 621 -PSTFDLSKWLFAQLNETIKQVHLSARPEFGEMGRITLYNFNVLADWYVMSKEVLY 679

5 Query: 662 SDNEDEKVTITSVLLATLTDQLRLHHPIMEFVTEIF--GQYAGSIVLASIYQVNAATFE 719
 D+E K R L Y LQILRLHHPMEFV +++ + SIV ASYF N FE
 Sbjct: 680 GDDEQAKAAKRVNLAYALDQLRLHHPMEFVHGKLMALHHTGKSIVTASYPVANTAFE 739

10 Query: 720 NQTHKGVESLKDILRSVPSNSRAEVNVAAPSKPITLVKTSDELSSESPKDNSTNIRPHT 779
 N A +++ LIR VR R E + ILVK +D L+ F+ N ++I RF N
 Sbjct: 740 NADATSAMDAILIINGVGRIGKAGAPLTKVDILVKITDPAKLPITFEQNPFDIFDRFVN 799

Query: 780 PETLEISSAATPELMASSSVITGAEIFLADLIANVEELARLEKELAKQKELDMVQKK 839
 + + + +A P+AA S+VITGA IF+FL +L++++EE A+L K+ K +E+ + KK
 Sbjct: 800 SKAFTVGTUVAEPKAGSAVITGATIFVPLNELIDLDEKAKCITDAKKEBQEIARDKK 859

15 Query: 840 LSNRPFVNAKPEVQKQKDEQTDYQTKDATTARIEMKK 880
 L+N+ F++ A VV +++ K+D++ + T R+E++++
 Sbjct: 860 LAKQGLSTAFEAUVAEQRTKRSDFDQLTSTKRLBQLR 900

20

A related DNA sequence was identified in *S.pyogenes* <SEQ ID 723> which encodes the amino acid sequence <SEQ ID 724>. Analysis of this protein sequence reveals the following:

Possible site: 61
 >>> Seems to have no N-terminal signal sequence

25 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.5062 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

30

An alignment of the GAS and GBS proteins is shown below:

Identities = 782/878 (89%), Positives = 818/878 (93%)

35 Query: 4 ELSPKYNPAEVEBGRQYCIWLDQVFKRSGTEAKPYSIVIPPNVTGKLHLGHAMUTTLQ 63
 ELSPKYNPAEVE GRQY WLD DVKPSGD +AKPYSIVIPPNVTGKLHLGHAMUTTLQ
 Sbjct: 3 ELSPKYNPAEVEAGRQYKWLDADVFKRSGDQAKPYSIVIPPNVTGKLHLGHAMUTTLQ 62

Query: 64 DIIIRQKRMQGFDTLKLPGMDHAGIATQAKVEERLRQGISRYDLGRDFLQKVVWEKDE 123
 DIIIRQKRMQGFDTLKLPGMDHAGIATQAKVEERLRQGISRYDLGR+KFLQKVVWEKDE
 40 Sbjct: 63 DIIIRQKRMQGFDTLKLPGMDHAGIATQAKVEERLRQGISRYDLGRDFLQKVVWEKDE 122

Query: 124 YAATIKESONGKGLSVDSYRERFTLDEGLSKAVRKVPVDLKNKGWIYRGFTINWDPAR 183
 YA TIK QWKGKGLSVDSYRERFTLDEGLSKAVRKVPVDLY KGMYRGFTINWDPAR
 45 Sbjct: 123 YATTIKESONGKGLSVDSYRERFTLDEGLSKAVRKVPVDLYKKGWIYRGFTINWDPAR 182

Query: 184 TALSDIEVIHKDVBGAFYHNNYMEDEGSRALVATTPETMFGDVAVAVNPEDARYKDLI 243
 TALSDIEVIHKDVBGAFYHNNYMEDEGSRAL+VATTPETMFGDVAVAVNPED RYKDLI
 50 Sbjct: 183 TALSDIEVIHKDVBGAFYHNNYMEDEGSRALQVATTPETMFGDVAVAVNPEDARYKDLI 242

Query: 244 GQNVILPIINKPIPIVADEHADPEFGTGVVKITPAHDPNDFAVGQRHNLQVNVMDGDT 303
 G+NVILPI+ NK IPIV DEHADPEFGTGVVKITPAHDPNDF VQRHNLQVNVMDGDT
 55 Sbjct: 243 GQNVILPIVNLPIPIVDEHADPEFGTGVVKITPAHDPNDFVQRHNLQVNVMDGDT 302

Query: 304 MNELADEFGMDRFEARKAVAKLESGLNVLKIKKTHSVGHISERTGVVPERLSTQWVF 363
 MNELA +F GMURFEAR+A VAKLE LG LV I+K HSGHISER+G VVPERLSTQWVF
 60 Sbjct: 303 MNELAGDPACMDRFEARQATVAKLEKGLVNLVTEKRVHSGHISERSGAVPERLSTQWVF 362

Query: 364 KMDQLAKNAJIANQJTDKVEFYPPRNDFTSMWNVHNDWISQRLWNGHQIPAWNVNG 423
 KMD+LAK A+ NQ+T+D+V+FYPPRNDFT+ WNVNVHNDWISQRLWNGHQIPAWN+G
 65 Sbjct: 363 KMDLEAKQAMNDGCTDTRVDFYPPRNDFTLQWNVNVHNDWISQRLWNGHQIPAWNVNG 422

Query: 424 EMYVGEDAPEGUQWTDQEDVLTWFSALWPFSTMGWNPDEAADPKRYPTISTLVGYDI 483
 E+YVGR+APEGD WTDQEDVLTWFSALWPFSTMGWNPOT+ DPKRYPTISTLVGYDI
 Sbjct: 423 EIVYGEAPEGUQWTDQEDVLTWFSALWPFSTMGWNPEDVDPKRYPTISTLVGYDI 482

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Query: 484 IFFVSRMIPQSLEPTGRQPPNVLIHGLIRDEBGRKMKSLGNGIDPMDVIEKYGDAL 543
 IFFVSRMIPQSLEPTGRQPP NVLIHGLIRDEBGRKMKSLGNGIDPMDVIEKYGDAL+L
 Sbjct: 483 IFFVSRMIPQSLEPTGRQPPNVLIHGLIRDEBGRKMKSLGNGIDPMDVIEKYGDAL 542

5 Query: 544 RWFLENGSAPQGDVRFPSYEHMDASWNFINKIWNISRYILMNEGLITLDAQARENVEKVVNS 603
 RWFLENGSAPQGDVRFPSYEHMDASWNFINKIWNISRYILMNEGLITL+ A NV KV S
 Sbjct: 543 RWFLENGSAPQGDVRFPSYEHMDASWNFINKIWNISRYILMNEGLITLEDAESNVAKVAAS 602

10 Query: 604 QVGNVTRWILHNLNETVGVKVTENFDKPERGVAGHILYNFIWEEFANWYVELTKEVLYSD 663
 + GNVTDWILHNLNET+ KVTENFDKPERGVAGHILYNFIWEEFANWYVELTKEVLYSD
 Sbjct: 603 SAGNVTDQWILHNLNETIAKVTENFDKPERGVAGHILYNFIWEEFANWYVELTKEVLYSD 662

15 Query: 664 NEDEKVTITRVSLLYTLDTQILRLHPIMPFVTERI+QYA+GSIV YP V FEN+ A 723
 NE EKVTITRVSLLYTLDT+ILRLHPIMPFVTERI+ QYA+GSIV YP V FEN+ A
 Sbjct: 663 NEAEKVTITRVSLLYTLDKILRLHPIMPFVTERIYAQYAGSIVTVDPVVRPAFENEA 722

20 Query: 724 HKGVESLKDILIRSVNSPRAEVNVAPSKPITILVKTSDESESFVKINSNYIKRFINPETL 783
 HKGVESLKDILIR+VRN+RAEVNVAPSKPITILVKT-DSELE FF N NYIK FINPE L
 Sbjct: 723 HKGVESLKDILIRAVNARAENVAPSKPITILVKTADSELEDFNSNINITYIKCFINPKL 782

Query: 784 EISSAIAPELAMSVITGAELPLADLINVEELARLEKELAKWKELDVMVGKLLGNE 843
 EISSAIA PELAM+S+ITGAEL+PLADLINVEELARL+KELAKWKELDVMVGKLL NE
 Sbjct: 783 EISSAIAPELAMSVITGAELPLADLINVEELARLEKELAKWKELDVMVGKLLGNE 842

25 Query: 844 RFVANAKPEVVQKEKDKQDTYQYKDATIARIEEMKKL 881
 RFVANAKPEVVQKEKDKQ DYQ KYDAT RI EMKK+
 Sbjct: 843 RFVANAKPEVVQKEKDKQADYQAKYDATQERIAEMKKI 880

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 228

A DNA sequence (GBSx0242) was identified in *S. agalactiae* <SEQ ID 725> which encodes the amino acid sequence <SEQ ID 726>. Analysis of this protein sequence reveals the following:

Possible site: 30
 >>> Seems to have no N-terminal signal sequence

----- Final Results -----
 bacterial cytoplasm --- Certainty=0.0669 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 727> which encodes the amino acid sequence <SEQ ID 728>. Analysis of this protein sequence reveals the following:

Possible site: 57
 >>> Seems to have a cleavable N-term signal seq.

----- Final Results -----
 bacterial outside --- Certainty=0.3000 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial cytoplasm --- Certainty=0.0000 (Not Clear) < succ>

The protein has no significant homology with any sequences in the GENPEPT database.

An alignment of the GAS and GBS proteins is shown below:

Identities = 148/191 (77%), Positives = 165/191 (85%)

Query: 14 GEKKKMNIIIGAAQSGKMTIGQIAKCTGMLFHHNDSIDFVLRFPWSPDSIALTESI 73

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G + KNN+LIIGAQAQSGKMTIGQE+A+QTGMTLPHNHSIDFVLRMPFWS +S AL E I
 Sbjct: 3 GAETYMNNLIIGAQAQSGKMTIGQEVARQTGMTLPHNHSIDFVLRMPFWSQSTALIERI 62
 Query: 74 RPKFFETFAKTGQGMILPTIVIDFNDSDRVVLEKIQIVFQSHNQEVLFVLETELSERLK 133
 RF FSTFAKTGQ+MEPTIVIDFND DV LKIQ VFGS++QEVLFVLEL+T++ SRLK
 Sbjct: 63 RFAFFETFAKTGQGMILPTIVIDFNDPNDVAMLEKIQAIVFQSYDQEVLFVLEKTDIERLK 122
 Query: 134 RNRTENRLKHKPSKRDIDNSESDDICSTMDYALFNPEVAPEALTYHKKINNTCLTATETAY 193
 RNRTENRLKHKP KR+I+MSE DI STM YA+FNPE P+ LT+Y KINNT LDA ETA
 Sbjct: 123 RNRTENRLKHKLGRILNLSBQDIQSTMAZAVFNPEEPKTLTHYKINNTQLTAATAAQ 182
 Query: 194 LIIQKINQIKE 204
 LIIQK+ IKE
 Sbjct: 183 LIIQKMIHIKE 193

Based on this analysis, it was predicted that these proteins and their epitopes could be useful antigens for vaccines or diagnostics.

Example 229

A DNA sequence (GBSx0243) was identified in *S. galactiae* <SEQ ID 729> which encodes the amino acid sequence <SEQ ID 730>. Analysis of this protein sequence reveals the following:

Possible site: 49
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.3614 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

The protein has homology with the following sequences in the GENPEPT database:

>GP:BAB04556 GB:AP001510 unknown conserved protein [Bacillus halodurans]
 Identities = 60/189 (31%), Positives = 102/189 (53%), Gaps = 3/189 (1%)
 Query: 7 EIVDNQLFVVETNRLLLRQKLEDAKEI FEFVGLDEVSYPAQFPFAVKSLEBETIYIETIY 66
 E + LP +ET RL LR+ +DA I+++ ++V+ + +S+++ ++ +
 Sbjct: 4 EDIYGDLPTELETERLRLLKFPYKDDAAAIYDASNEQVTKYVLWETHQSIIKDSRAFLA--F 61
 Query: 67 PTNLEKEKLPSGYAITLNGDDKVIQSVDFNH-RHEDDIFETGYLLHPYWGQGI+VPEAS 125
 N EK S +AL LK +++IG+VDF + +D E+GY+L YWQGQI+ EA +
 Sbjct: 62 ALNKYDEKDVSPWALBGRNRMIGTVDFVWVKFDRKTARLGYLVSEPYWGQGITAVN 121
 Query: 126 ALVEIGFTLLGLHKIELGCVYDNKQSQAVARKLGFTRANIRDRRDAGGKROGDMRGLL 185
 ALVE GF + L +L+ C+ N S V K G E R +G + ++
 Sbjct: 123 ALVEFGFNMELEKRIQAUCFARNISSARVMSKAGLIYTBGTHRRATYVKGARHDFKYAYII 181
 Query: 186 RSEWEKKRR 194
 R ++E+K +
 Sbjct: 182 REDYEQKHQ 190

A related DNA sequence was identified in *S. pyogenes* <SEQ ID 731> which encodes the amino acid sequence <SEQ ID 732>. Analysis of this protein sequence reveals the following:

Possible site: 30
 >>> Seems to have no N-terminal signal sequence
 ----- Final Results -----
 bacterial cytoplasm --- Certainty=0.1864 (Affirmative) < succ>
 bacterial membrane --- Certainty=0.0000 (Not Clear) < succ>
 bacterial outside --- Certainty=0.0000 (Not Clear) < succ>

An alignment of the GAS and GBS proteins is shown below: